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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:

C12N 15/31, 15/62, 1/21, C07K 14/245, C12Q 1/00

(11) International Publication Number:

WO 95/20657

(43) International Publication Date:

3 August 1995 (03.08.95)

(21) International Application Number:

PCT/DK95/00042

A1

(22) International Filing Date:

27 January 1995 (27.01.95)

(30) Priority Data:

08/187,166

27 January 1994 (27.01.94) US

(60) Parent Application or Grant

(63) Related by Continuation

US Filed on 08/187,166 (CIP) 27 January 1994 (27.01.94)

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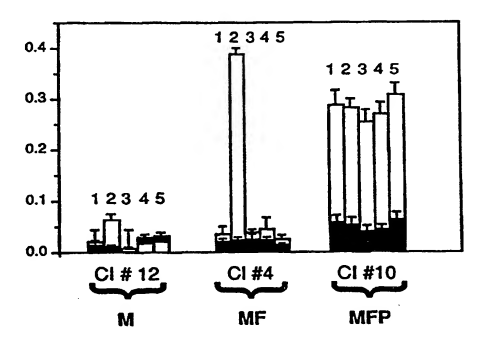
(81) Designated States: AM, AT (Utility model), AU, BB, BG, BR, BY, CA, CN, CZ, CZ (Utility model), DE (Utility model), DK (Utility model), EE, ES (Utility model), FI, FI (Utility model), GE, HU, JP, KG, KP, KR, KZ, LK, LR, LT, LV, MD, MG, MN, MX, NO, NZ, PL, RO, RU, SI, SK, SK (Utility model), TJ, TT, UA, US, UZ, VN, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG), ARIPO patent (KE, MW, SD, SZ).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: RECEPTOR SPECIFIC BACTERIAL ADHESINS AND THEIR USE



CLINICAL ISOLATES

(57) Abstract

Bacterial adhesins that have been selected or recombined to have the ability to bind specifically to pre-determined, selected inanimate or animate receptors and the use of such adhesins or bacteria expressing the adhesins, in the targeting of useful compounds and/or bacteria to selected cells and surfaces.

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RECEPTOR SPECIFIC BACTERIAL ADHESINS AND THEIR USE

FIELD OF INVENTION

The present invention pertains to naturally occurring bacterial adhesins and derivatives and variants hereof, having the ability to bind to pre-determined, specifically selected receptors, and to the use of such adhesins in the targeting of active compounds and microbial cells to locations comprising such selected receptors.

This invention was supported in part by the US National

10 Institute of Health (NIH), under grant #DE07218, and the US

Veterans Administration. The US government has certain rights
in the invention.

TECHNICAL BACKGROUND AND PRIOR ART

The ability to adhere or bind specifically to, and in many instances, to colonize an animate or inanimate surface is of paramount importance in microbial ecology and pathogenesis. Such specific receptor binding is provided by microbial adhesins which play a key role in bacterial/host and viral/host recognition and interaction and for the recognition of any specific surface by a microorganism.

Accordingly, adhesion of bacteria to host surfaces is commonly regarded as an essential step enabling bacteria to become established as members of the normal flora of host organisms or to cause an infection (refs. 7, 18). Bacterial lectins are the most common and most thoroughly studied type of adhesins among both gram negative and gram positive bacteria (ref. 40). Evolutionary pressures have selected lectins for adhesive functions probably due to the abundance of glycoconjugates on animate and inanimate surfaces. One class of structures that a large range of gram-positive and gram-positive bacteria including Escherichia coli and other members of the

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family Enterobacteriaceae, have evolved to adhere to host glycoproteins in a saccharide-dependent manner are surface fibrils called fimbriae (ref. 14) or pili (ref. 10). Colonization Factor Antigen (CFA) type I and Colonization Factor Antigen (CFA) type II are specific examples of such fimbriae.

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By far the most common of the enterobacterial fimbriae is type 1, or mannose-specific (MS) fimbriae (refs. 11, 13, 14, 23). Type 1 fimbriae are heteropolymers of four different subunits (refs. 28, 44). For each fimbria, about 1000 copies of a 17-kDa primary structural subunit designated FimA (or PilA), are polymerized into a right-handed helix surrounding a hollow axial core (ref. 11). Three ancillary subunits, FimF, FimG and FimH, are also polymerized into the fimbrial structure, but comprise only 1-3% of the fimbrial mass (refs. 20, 24, 27, 32).

The 28 kDa FimH subunit has been shown by several direct and indirect tests to be the actual fimbrial lectin (refs. 2, 4, 20, 21, 27, 29, 32, 36, 55), although its function may be affected by other subunits (ref. 55). The FimA subunit is highly variable, but the FimH subunit is highly conserved antigenically and genetically among enterobacteria (ref. 1). Interactions between type 1 fimbriae and D-mannose-containing receptors have been shown in a number of studies to play a key role in the infectious process (refs. 2, 4, 9, 19, 25, 26, 31, 33, 44, 50).

Detailed analysis of adhesion-inhibition or agglutination-inhibition by various mannosides and manno-oligosaccharides have suggested that the combining site of the type 1 adhesin is in the form of an extended pocket corresponding to the size of a trisaccharide and fitting best the structure α -D-Manp-(1-3)- β -D-Manp-(1-4)-D-GlcNac (ref. 16). A hydrophobic region within or close to the combining site was also predicted in these studies. A similar pattern of specificity was found independently in indirect adhesion-inhibition studies, as well as in direct adhesion studies using "neoglycolipids"

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as receptors (refs. 37, 47). The combining site of the Klebsiella pneumoniae type 1 adhesin was shown to be similar to the Escherichia coli adhesin, whereas the Salmonella typhimurium type 1 adhesin combining site appears to be smaller and devoid of a hydrophobic region (ref. 16) Thus, it has long been thought that type 1 fimbriae of enterobacteria were functionally quite similar and that the primary essential characteristic of any potential receptor was the presence of terminal α 1-3-linked mannosyl residues.

- Recently it has been reported that the type 1 fimbriated, K-12-derived E. coli strain CSH-50 exhibits mannose-sensitive peptide-binding activity (ref. 51). CSH-50 E. coli bound to yeast mannan (Mn), a highly mannosylated glycoprotein, and to human plasma fibronectin (Fn) when immobilized on assay
- wells. Adhesion to Mm, but not to Fn, was essentially eliminated by periodate treatment. Furthermore, CSH-50 E. coliadhered in a mannose-sensitive fashion to non-glycosylated peptide fragments of Fn and to a synthetic peptide copying the first 30 residues of the Fn molecule, FnSp1. Fimbriae
- purified from these organisms also bound to Fn and FnSp1. A well-characterized recombinant strain of E. coli PC31 expressing type 1 fimbriae, HB101(pPKL4), adhered to Mn, but did not adhere to the other substrata. Fimbriae purified from HB101(pPKL4) did not adhere to Fn or FnSp1. Thus, E. coli
- 25 type 1 fimbriae appeared to be functionally heterogeneous.

Several *E. coli* isolates obtained from human urine also expressed peptide-binding activity similar to that of CSH-50, indicating that this new phenotype was not restricted to a laboratory strain. Other isolates expressed an adhesive activity similar to that of HB101(pPKL4). A third class of type 1 fimbriae-mediated adhesive phenotype was also observed among these isolates.

The FimH subunit is the D-mannose-sensitive adhesin of type 1 fimbriae, common i.a. to the *Enterobacteriaceae*. It is presently widely accepted that host receptors are strictly

limited to glycoproteins containing terminal mannosyl residues (refs. 16, 37, 41, 42, 43, 47). Hereinbelow functional and genetic evidence is provided demonstrating that this generalization is not correct. Allelic variants of *E. coli* fimH genes encoding proteins differing by as little as a single amino acid substitution confer distinct adhesive phenotypes and accordingly, the fimH gene is not a single gene but rather a family of fimH genes.

Surprisingly, active receptors for FimH proteins were found to include glycoprotein domains where mannosyl residues are not terminal and even protein domains devoid of saccharide. This unexpected adhesive diversity within the fimH family broadens the scope of potential receptors for bacterial adhesion and may lead to a fundamental change in the understanding of the role(s) type 1 fimbriae and other bacterial adhesins may play in bacterial ecology or pathogenesis.

The present findings also opens up a completely new field of technology, since it provides the means to design bacteria expressing adhesins that bind to pre-determined, specific receptors in a wide range of animate and inanimate locations. This new technology is referred to herein as Designer Adhesin Technology.

SUMMARY OF THE INVENTION

Accordingly, the present invention relates in one aspect to a recombinant or mutant bacterial adhesin variant derived from a naturally occurring adhesin, said adhesin variant having altered binding properties relative to the naturally occurring adhesin from which it is derived.

In further aspects the invention provides a FimH adhesin

30 having an amino acid sequence which differs from the *E. coli*PC31 FimH adhesin by at least one amino acid and a recombinant replicon comprising a DNA sequence selected from the

group consisting of a sequence coding for a recombinant bacterial adhesin variant as defined above and a sequence coding for a FimH adhesin as also defined above.

In a still further aspect, there is provided a fusion protein comprising an adhesin selected from the group consisting of a recombinant bacterial adhesin variant as defined above and a FimH adhesin as also defined above, and a heterologous polypeptide.

The invention also pertains to a recombinant bacterial cell
which expresses an adhesin selected from the group consisting
of a recombinant bacterial adhesin variant as defined above
and a FimH adhesin as defined above, and to a composition
comprising a population of such cells.

In one interesting aspect of the invention there is provided $ilde{\sim}$ 15 a method of isolating a bacterial cell expressing an adhesin having modified binding properties relative to a natively 1 expressed adhesin, comprising identifying in the bacterial cell DNA sequence(s) coding for the binding domain(s) of said natively expressed adhesin and substituting at least one 20 codon herein, whereby a modified adhesin molecule is expressed that is different in at least one amino acid from the adhesin expressed natively, and selecting a bacterial cell expressing the modified adhesin having an altered adhesion phenotype relative the natively expressed bacterial 25 adhesin.

In a further interesting aspect the invention relates to a method of preparing a recombinant bacterial cell that binds to a specific receptor moiety, comprising introducing into a bacterium that does not produce an adhesin binding to said receptor moiety, a DNA sequence coding for an adhesin binding to the receptor moiety, and selecting a bacterial cell expressing the DNA sequence.

There is also provided a method of targeting a bacterial adhesin to a specific location, comprising (i) identifying in said location an adhesin-interacting receptor moiety which is recognizable by bacterial adhesins, said moiety preferably being one which is occurring abundantly, (ii) isolating a bacterial cell that grows in said location and expresses an adhesin recognizing and interacting with said receptor moiety, and administering to the location the bacterial cell or the adhesin under conditions where the adhesin and the receptor moiety are brought into interacting contact whereby the adhesin is associated with the receptor moiety.

DETAILED DISCLOSURE OF THE INVENTION

As used herein the term "bacterial adhesins" denotes proteins which recognize and bind to a large variety of target molecules such as polysaccharides, glycolipids, glycoproteins, polypeptides and proteins. More than a hundred different adhesins have been described so far originating from a large variety of gram-negative and gram-positive bacteria. Adhesins can be present on the bacterial surface as components of organelles such as fimbriae, also called pili or fibrillae, these three terms being used interchangeably herein, or as non-fimbrial or afimbrial adhesins (ref. 64). Examples of fimbrial or pili adhesins include the following surface structures in E. coli: P pili, type 1 fimbriae, S pili, K88 pili, K99 pili, CS3 pili, F17 pili and CS31 A; in Klebsiella pņeumoniae: type 3 pili; in Bordetella pertussis: type 2 pili; in Yersinia enterocolitica: Myf fibrillae; in Yersinia pestis: pH6 antigen and F1 envelope antigen.

Examples of non-fimbrial cell surface structures which have

adhesin function or which may comprise proteins having such a
function include capsules, lipopolysaccharide layers, outer
membrane proteins, NFA (non-fimbrial adhesin)-1, NFA-2, NFA
3, NFA-4, AFA (afimbrial adhesins)-I, AFA-II and AFA-III.

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In the present context, the term "fimbriae" designates long thread-like bacterial surface organelles. Fimbriae are heteropolymers each consisting of about 1000 structural components, mostly of a single protein species. However, in many cases a few percent minor components are also present. Adhesins can either be identical to the major structural protein as in Escherichia coli K88 and CFA1 fimbriae and type 4 fimbriae of Pseudomonas, Vibrio and Neisseria, or they may be present as minor components as in E. coli type 1 and P fimbriae [for reviews see Krogfelt 1991 (ref. 62); Kaufman 10 and Taylor, 1994 (ref. 60): Kuehn et al., 1994 (ref. 63); Klemm and Krogfelt, 1994 (ref. 61)]. In the latter case, i.e. when present as minor compounds, the adhesins are closely related in amino acid sequence to the major fimbrial component. As used herein the term bacterial adhesin will also 15 include adhesins isolated from non-bacterial sources including viruses, and which are expressed in a bacterium.

In the following, the FimH adhesin of type 1 fimbriae will be described structurally and functionally as a representative example of a bacterial adhesin.

FimH is located at the tip of the type 1 fimbriae and also intercalated at intervals in the fimbrial organelle. Most forms of the FimH adhesin target to (bind to) oligosaccharide structures containing terminally located α -D-mannoside residues [Krogfelt et al., 1990 (ref. 29)]. Based on studies with various D-mannose derivatives the receptor binding site of the FimH adhesin is assumed to be shaped like an elongated pocket large enough to accommodate a trisaccharide motif [Sharon, 1987 (ref. 65)].

The fimH gene encodes the precursor FimH protein of 300 amino acids [Klemm and Christiansen, 1987 (ref. 27)]. This precursor is processed into a mature form of 279 amino acids. The amino acid sequence of the E. coli PC31 FimH protein is shown in Table 1 below wherein cysteine residues are indicated by asterixes, the signal peptide is outlined in bold letters,

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and two regions contributing to the binding site are underlined (SEQ ID NO:1). (It should be noted that residue 176 is a proline residue and not as previously indicated when the PC31 FimH protein was first published, an arginine residue):

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Table 1. Amino acid sequence of the E. coli PC31 FimH protein

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 $\textbf{MKRVITLFAVLLMGWSVNAWS} \textbf{FACKTANGTAIPIGGGSANVYVNLAPVVNVG} \underline{\textbf{ONLVVDLS}}$

10 TOIFCHNDYPETITDYVTLORGSAYGGVLSNFSGTVKYSGSSYPFPTTSETPRVVYNSRT

DKPWPVALYLTPVSSAG<u>GVAI</u>KAGSLIAVLILRQTNNYNSDDFQFVWNIYANNDVVVPTG

GCDVSARDVTVTLPDYPGSVPIPLTVYCAKSQNLGYYLSGTHADAGNSIFTNTASFSPAQ

15 GVGVQLTRNGTIIPANNTVSLGAVGTSAVSLGLTANYARTGGQVTAGNVQSIIGVTFVYQ

The FimH contains 4 cysteine residues assumed to direct folding of the molecule into distinct functional domains. For comparison FimA and the minor components FimF and FimG only have two cysteine residues. The localization of the cysteine residues in FimH points to a tandem arrangement of two ancestral genes. Furthermore, similar amino acids can be found in similar positions in the two halves of the FimH protein. The "midway" point is located roughly around residue 150 in the mature protein. The two halves or domains of FimH have evolved differently with the N-terminal section becoming the domain harbouring the receptor binding site, whereas the C-terminal sector became the domain of the molecule required for integration into the fimbrial organelle structure, i.e. having the features of a structural component.

In-frame linker insertions into the fimH gene confirms this model of the FimH protein. Thus insertions in the C-terminal half of the molecule generally do not interfere with the

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receptor-binding ability whereas abolishment of receptor binding ability following linker insertion in the N-terminal is the rule (Klemm et al., unpublished data). A similar domain structure has been observed in the PapG adhesin of P-fimbriae [Hultgren et al., 1989 (ref. 59); Kuehn et al., 1994 (ref. 63)].

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In accordance with the invention, the recombinant bacterial adhesin as defined above is one which is derived from an adhesin having certain binding properties, but which recombinant bacterial adhesin has altered binding properties relative to the naturally occurring adhesin (the parent adhesin) from which it is derived. As used herein this feature encompasses situations where the adhesin variant recognizes and binds to receptor moieties not being recognized by the parent adhesin irrespective of whether the adhesin variant has lost its normal ability to recognize and bind to a certain receptor moieties, or not.

As used herein the term "binding" indicates that the adhesin has a degree of affinity to the receptor moiety which enables it, when brought into contact herewith, to interact in a binding manner with this moiety whereby an adhesin-receptor moiety association occurs. The strength of this binding depends on the type of binding force which causes the interaction between the receptor moiety and the adhesin. In the present context, such binding forces include covalent binding and binding by non-covalent binding forces including hydrogen bonds, hydrophobic interactions, van der Waal forces and ionic interactions. Accordingly, the term "receptor moiety" as used herein encompasses any moiety to which an adhesin may interact by the above binding forces.

In one specific embodiment, the adhesin variant is a FimH mannose-sensitive adhesin normally binding to a receptor selected from a domain where mannosyl residues are not terminal and a domain devoid of saccharide and having an amino acid sequence which differs from the *E. coli* PC31 FimH adhes-

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in by at least one amino acid residue substitution, including an amino acid sequence differing by at least 2 amino acids, preferably by at least 3 amino acids, more preferably by at least 4 amino acids, most preferably by at least 5 amino acids. In further useful embodiments, the amino acid sequence may even differ by more than 5 amino acids such as at least 6, preferably by at least 7, more preferably by at least 8, even more preferably by at least 9 and in particular by at least 10 amino acid residues, such as by at least 12 amino acids including by at least 15.

Accordingly, the above FimH adhesin variant is preferably at least 90% homologous to the PC31 FimH adhesin, such as at least 92% homologous, more preferably at least 93% homologous, even more preferably at least 94% homologous, most preferably at least 95% homologous, and in particular at least 96% homologous, e.g. at least 97% homologous. In particularly interesting embodiments, the adhesin is at least 98% homologous, including at least 99% homologous such as at least 99.5% homologous.

The above FimH adhesin variant can be a chimeric adhesin comprising amino acid sequences from different FimH adhesins having identical or different binding specificities.

As it has been mentioned above, the present invention is generally aimed at providing the means to design bacterial adhesins having specific binding properties whereby bacteria expressing the adhesin variants or the adhesin variants in isolated or purified form can be designed to bind to a specific desired target receptor moiety. Accordingly, the adhesin variant may in accordance with the invention be an adhesin variant as defined above which binds to an animate receptor moiety. Such receptors include receptors located on inner surfaces of humans and animals, such as e.g. the mucosal membranes of the gastrointestinal tract including the teeth and the oral cavity, and the mucosal membranes of the respiratory and the genitourinary systems. Included are also

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adhesin variants that bind to outer surfaces, including the skin, of humans and animals.

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In a further embodiment, the adhesin variant is designed so as to acquire the ability to bind to a plant receptor moiety. This aspect is of particular interest in relation to deliber-5 ate release to out-door or in-door environments where plants are cultivated, of useful recombinant bacteria having a desirable effect on the growth and yield of the plants. Such desirable bacteria are e.g. bacteria expressing a pesticidally active substance, i.e. a biopesticide including as examples pesticidal toxins produced naturally by Bacillus spp such as the Bacillus thuringiensis (Bt) toxin. In this context, another example is bacteria which protect plants against low temperature damages or bacteria which express gene products protecting plants against detrimental effects of herbicides.

By providing such bacteria with genes expressing adhesin variants which e.g. bind specifically to certain plant species and/or to certain locations on the plant, these useful bacteria will, when administered to the plant growing environment, be selectively associated with the target plant species or a specific target area of the plant. It may thus be desirable to have these useful bacteria administered to the leaves of the plants or to have the root system colonized herewith.

Accordingly, the present invention encompasses adhesin variants as defined herein which bind selectively or specifically to a phylloplane receptor moiety or which bind to receptors on plant roots. Similarly, adhesin variants can be provided which are targeted to the stem or the flowers of the plants.

As it is mentioned above, bacterial adhesins include adhesins having an inherent capability to bind or interact with inanimate surfaces carrying receptor moieties with which the adhesin can interact to become bound to the surfaces. It is

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known that certain bacterial adhesins can bind to inanimate surfaces including as examples glass, hydroxyapatite (a tooth enamel model compound) or polymer structures including plastics and polysilicates. The present invention has made it possible to design bacteria which bind selectively to any inanimate surface which carries a receptor moiety for which an adhesin variant binding thereto may be constructed. Accordingly, the present invention also provides an adhesin variant as defined herein which binds to an inanimate receptor 10 moiety. Such adhesin variants are particularly interesting in connection with the concept of bioremediation, i.e. a technology designed to enhance degradation of chemical pollutants in the environment. It is clearly a significant improvement of this technology to have at hand bacteria which comprise genes coding for pollutant-degrading gene products and which also express adhesins targeting the bacteria selectively to the environment where the pollutants are present, e.g soil, aquatic environments and drinking water supply systems. Furthermore, adhesin variants capable of binding to tooth enamel are useful in the protection of teeth against caries.

In a further embodiment, there is in accordance with the invention provided an adhesin variant which is part of a fusion protein comprising the adhesin variant and a nonadhesin, heterologous polypeptide. Using the FimH as an example, it has been found that fusions between a bacterial adhesin and other proteins can be made whereby the resulting fusion proteins are inserted into the cell surface organelle of which the adhesin is a structural part. These resulting hybrid adhesin-carrying cell organelles remain fully functional with respect to binding properties. Additionally, it has been found that large regions of non-adhesin proteins, e.g. regions comprising in the range of 1 to 100 amino acids including a range of 5 to 75 amino acids and a range of 10 to 60 amino acids, such as regions comprising 15 to 54 amino acids, can be inserted into type 1 fimbriae without impairing the binding properties of the fimbriae.

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In useful embodiments of the invention, the non-adhesin region of a fusion protein comprising an adhesin variant as defined herein include a heterologous polypeptide which is selected from an epitope, an enzyme, a toxic gene product and an antibody.

It has significantly been found that, when fusion proteins are expressed in which the heterologous polypeptide is an epitope or an epitope-carrying domain forming an integrated part of the fusion protein, and thus presented on the surface of the host cell expressing the fusion protein, the epitope-carrying polypeptide can be presented in a conformational form similar to its natural conformation.

Furthermore, it has surprisingly been found that the above fusion proteins can be overproduced by the bacteria comprising hybrid genes coding for fusion proteins, resulting in excretion of the fusion proteins to the growth medium in large quantities. Accordingly, the excreted fusion proteins are then readily isolated and purified, e.g. by means of affinity chromatography. These findings provide the means to manufacture bacterial cells having on their surface hybrid adhesin-carrying cell organelles as well as to produce large quantities of excreted fusion proteins, both of which can be targeted to specific surfaces as determined by the binding properties of the adhesin variant of the fusion protein.

- The above technology of making adhesin variant-fusion proteins is useful for a range of industrially important purposes such as:
 - (i) development of live vaccines targeted to specific cellular surfaces;
- (ii) development of subunit vaccines for administration orally or by injection, which are targeted to pre-determined, specifically selected cell surfaces or mucosal surfaces;

(iii) development of fusion proteins combining specific binding properties with specific enzymatic or toxin activities. Such fusion proteins have applications as therapeutical or diagnostical agents, including use in biosensors;

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- (iv) use of fusion proteins as carriers of non-covalently linked chemical moieties whereby the adhesin part of the protein is used to target the chemical moiety to specific locations and the non-adhesin part carries and then releases the moiety when the fusion protein has reached its target.
- Examples of chemical entities which may be linked to the 10 fusion protein include imaging agents and pharmacologically active components. Examples of applications for this use include imaging of atherosclerotic plaques or tumor tissues, and delivery of chemical agents at specific locations in or on microbial, human, animal or plant cells including specific 15 tissues or tissue components;
 - (v) development of fusion proteins which are useful in affinity purification processes.

It has been found that the fimH gene coding for the $E.\ coli$ FimH adhesin is not a single gene but rather a family of fimH 20 genes, and accordingly it has now been established that allelic variants of E. coli fimH genes exist that encode adhesin proteins which, relative to the known $E.\ coli$ PC31 fimH gene product differ by as little as a single amino acid substitution and confer distinct binding or adhesive 25 phenotypes.

Accordingly, as it has been mentioned above, the present invention relates in a further aspect to a FimH adhesin having an amino acid sequence which differs from the E. coli PC31 FimH adhesin as defined above by substitution of at 30 least one amino acid. It will be understood that such an adhesin encompasses naturally occurring adhesins as well as adhesins which are encoded by recombinant or mutant fimH genes. In this context the term "fimH gene" denotes a gene

coding for a gene product which can be integrated into a type 1 fimbria and which confers to the fimbria the ability to recognize and bind to a receptor.

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The FimH adhesin as defined above may be an adhesin having its inherent binding properties or an adhesin variant which in relation to an adhesin encoded by a naturally occurring gene from which the gene coding for the adhesin variant is derived, has altered binding properties. Furthermore, the FimH adhesin may be either mannose-sensitive or mannoseinsensitive. The term "mannose-sensitive" is used herein to 10 designate that the binding of an adhesin is inhibited in the presence of mannose residues. In one specific embodiment, the FimH adhesin may be a FimH adhesin normally binding to a receptor moiety selected from a domain where mannosyl residues are not terminal and a domain devoid of saccharide such 15 as e.g. a glycolipid, a glycoprotein, a protein, a polypeptide and a peptide, including a hormone. Examples of proteins to which a FimH adhesin according to the present invention may bind include as examples animal proteins such as a casein including κ -casein, a gelatine, a globin, an 20 albumen and a collagen, and vegetable proteins including soy protein.

The FimH adhesin according to the invention include an adhesin having an amino acid sequence which differs from the E.

25 coli PC31 FimH adhesin by at least 2 amino acid residues, such as an amino acid sequence differing by at least 3 amino acids, preferably by at least 4 amino acids, more preferably by at least 5 amino acids, most preferably by at least 6 amino acids. In further useful embodiments, the amino acid sequence may even differ by more than 6 amino acids such as at least 7, preferably by at least 8, more preferably by at least 9, even more preferably by at least 10 and in particular by at least 11 amino acid residues, such as by at least 12 amino acids including by at least 15.

Accordingly, the above FimH adhesin is preferably at least 90% homologous to the PC31 FimH adhesin, such as at least 92% homologous, more preferably at least 93% homologous, even more preferably at least 94% homologous, most preferably at least 95% homologous, and in particular at least 96% homologous, e.g. at least 97% homologous. In particularly interesting embodiments, the adhesin is at least 98% homologous, including at least 99% homologous or at least 99.5% homologous.

In one specific embodiment, the FimH adhesin as defined above is one which, when tested for binding to yeast mannan (Mn), human plasma fibronectin (Fn), periodate treated Fn and the synthetic peptide FnSpl comprising the first 30 amino acids of Fn, only binds to Mn. In the following, an adhesin having this pattern of binding properties is designated an M class FimH adhesin. In other specific embodiments, the FimH adhesin is an adhesin which, when tested for binding to the above compounds, binds to Mn and Fn (MF class FimH adhesin) or an adhesin which among these compounds bind to all of these (MFP class FimH adhesin).

It has been found that bacteria expressing FimH adhesins of the above MFP class bind in a mannose-sensitive (MS) manner to polyoxyethylene sorbitan monolaurate (Tween 20) and a little less well to polyoxyethylene sorbitan monooleate 25 (Tween 80). Furthermore, bacteria expressing MFP class FimH adhesins make a much tougher pellicle than bacteria expressing other types of adhesins. In the present context, the term "pellicle" indicates a layer or film of bacteria that forms at the air/liquid interface of a liquid growth medium. This noticeable phenomenon might be of particular interest 30 where there is a reason to concentrate microorganisms at the surface of an aquatic environment, such as e.g. bacterial cells which in accordance with the present invention express a pollutant-degrading gene product.

Another interesting finding is that bacteria expressing a MFP class adhesins bind to hydroxyapatite to a higher degree than do bacteria expressing a M class adhesin. Hydroxyapatite, especially saliva-treated hydroxyapatite is i.a. used as a model for tooth enamel, and accordingly, this finding indicates that bacteria expressing MFP class adhesins are particularly useful in bacterial compositions intended for colonization of teeth.

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It has also been found that the MFP class adhesins bind to a large range of synthetic peptides and accordingly seem to 10 have a broad specificity in terms of amino acid motifs.

In further specific embodiments of the invention, the FimH adhesin is an adhesin which, when tested for binding to the five Fn-fragments obtained by thermolysin treatment as it is described in reference No. 51, only binds to the 40-kDa gelatin-binding fragment or which binds to all of these Fnfragments, or to none of these.

In addition to the above classes of FimH adhesins, another class has been identified which is designated the M^{L} (low 20 adhesive) class. Such an adhesin confers the ability to aggregate yeast cells in a mannose-sensitive (MS) fashion, in titers similar to M class adhesins, but surprisingly, it binds at only low levels to Mn or Fn and FnSp1. Furthermore, adhesins of this low adhesive M^{L} class adhere poorly to MDCK, 25 buccal cells and erythrocytes as compared with M class adhesins. Example of a M^L class adhesin is one expressed by the recombinant E. coli strain KB 23 which differs only from the PC31 FimH adhesin by having an alanine instead of a valine at residue 27 and the FimH adhesin expressed by the human fecal 30 E. coli isolate which is designated F-18 [McCormick et al., 1989 (ref. 34)]. This latter adhesin differs from the PC31 FimH in three amino acid residues and the F-18 isolate has been found to colonize the large intestine to a higher degree than certain E. coli K-12 strains do. Accordingly, it is contemplated that these ML class adhesins confer to

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gastrointestinal bacteria the ability to colonize the large intestine which is significant for a live bacterial vaccine for exerting its immunological effect in the gastrointestinal tract.

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Furthermore, it has been found that among M class adhesins adhesion is found that is not sensitive to inhibition by Dmannose. Such a mannose-insensitive (or mannose-resistant) ${\tt M}$ class adhesin is designated in the following as an M^R adhesin. One example of a bacterial strain expressing an M^R adhesin is the clinical isolate U221-3 which is mentioned in the 10 following.

In accordance with the invention, a FimH adhesin as defined above can be a chimeric adhesin comprising amino acid sequences from different FimH adhesins. Such chimeras are constructed e.g. by providing multiple restriction fragments 15 of a fimH gene, followed by exchanging under ligation conditions these fragments with corresponding fragments of an other fimH gene and cloning the ligation product as it is described in Example 1 below. As it is also explained below, recombinant plasmids containing such chimeric fimH genes can be transformed into a host cell and transformants tested for adhesive phenotype, allowing determination of the regions of each gene capable of conferring functional activity (Fig. 5). These studies which are described in details below showed that all of the sequence changes relative to the PC31 fimH gene that affected binding function in the studied strains of E. coli CSH-50 and clinical isolates (CIs) designated #s 3, 4, 7, 10, F-18 and U221-3, respectively, occurred between residues 27 and 119, both included, of the 279 residue, mature fimH sequences.

Accordingly, the invention encompasses in one embodiment a FimH adhesin comprising an amino acid sequence which differs from the $E.\ coli$ PC31 FimH adhesin by at least one amino acid occurring between residues 27 and 119 of the mature FimH sequence, including a FimH adhesin comprising an amino acid 35

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sequence which differs from the $E.\ coli$ PC31 FimH adhesin by at least one amino acid occurring between residues 33 and 78 of the mature FimH sequence.

The selected potential receptors for a FimH adhesin as defined above include those animate and inanimate receptors mentioned above for a recombinant bacterial adhesin variant and the potential uses of the FimH adhesins are also the same as those uses described above for this recombinant bacterial adhesin variant.

10 As mentioned above, the invention relates in a further aspect to a recombinant replicon comprising a DNA sequence coding for a recombinant bacterial adhesin variant as defined herein or a DNA sequence coding for a FimH adhesin as also defined herein. Such a replicon is suitably selected from a chromosome or a plasmid. The DNA sequence includes a sequence which 15 is inserted by conventional recombination techniques such as insertion by means of restriction enzymes and subsequent ligation, or the DNA sequence is provided by subjecting a replicon comprising a naturally occurring sequence coding for an adhesin to a mutagenization procedure including site-20 directed mutagenesis, insertion of a transposable element, mutagenization by radiation or chemical mutagenization, followed by selection of cells comprising a mutated sequence conferring altered binding properties relative to a cell 25 comprising the wild-type sequence.

In preferred embodiments, the recombinant replicon is one having a broad host range including bacterial species naturally occurring in soil, in aquatic environments, on inner and outer surfaces of humans and animals, and which is compatible with replicons occurring in potential host strains.

In one useful embodiment, the recombinant replicon as defined above is one wherein the DNA sequence codes for a FimH adhesin having an amino acid sequence which differs from the E. coli PC31 FimH adhesin by at least one amino acid, including

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an adhesin having an amino acid sequence which differs from the E. coli PC31 FimH adhesin by at least 2 amino acid residues, such as an amino acid sequence differing by at least 3 amino acids, preferably by at least 4 amino acids, more preferably by at least 5 amino acids, most preferably by at least 6 amino acids. In further useful embodiments, the amino acid sequence may even differ by more than 6 amino acids such at least 7, preferably by at least 8, more preferably by at least 9, even more preferably by at least 10 and in particular by at least 11 amino acid residues, such as by at least 12 amino acids including by at least 15.

Accordingly, the above recombinant replicon preferably comprises a DNA sequence coding for a FimH adhesin which is at least 90% homologous to the PC31 fimH gene, such as at least 92% homologous, more preferably at least 93% homologous, even 15 more preferably at least 94% homologous, most preferably at least 95% homologous, and in particular at least 96% homologous, e.g. at least 97% homologous. In particularly interesting embodiments, the adhesin is at least 98% homologous, including at least 99% homologous such as at 20 least 99.5% homologous.

In a further embodiment, the above replicon comprises a DNA sequence which is a chimeric fimH gene as it has been defined above, comprising DNA from different fimH genes. The replicon can also be one which comprises a further DNA sequence e.g. 25 derived from a microorganism selected from a bacterium, a virus, a protozoan, a fungus and a yeast. This further DNA sequence is e.g. one coding for a heterologous polypeptide, including an epitope, an antibody, a toxic gene product, an enzyme, a pesticidally active gene product and a pollutantdegrading gene product.

In useful embodiments, the replicon as defined herein comprises a DNA sequence which is isolated from an Enterobacteriaceae species, including a DNA sequence which is isolated

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from E. coli, a Klebsiella sp., an Enterobacter sp., a Yersinia sp. or a Salmonella sp.

In addition to being a DNA sequence as defined above, the sequence can be a synthetic sequence constructed by conventional techniques of DNA synthesis.

As it is also mentioned above, the present invention encompasses a fusion protein comprising a recombinant bacterial adhesin variant or a FimH adhesin as defined above, and a heterologous polypeptide. Such a polypeptide is in useful embodiments an immunologically active gene product i.e. an 10 epitope (antigenic determinant) from a pathogenic organism, which polypeptide, when administered to the body of a human or an animal is capable of stimulating the formation of antibodies therein. A cell in which such an epitope is expressed is advantageously utilized in the preparation of live vaccines. Such vaccines have several advantages over known live vaccines:

Firstly, the epitope forms a structural part of an adhesin which is embedded in a surface organelle of the vaccine cells. This implies that the hybrid DNA sequence coding for 20 the epitope further comprises the means for transporting the epitope, when expressed, to the outer surface of the cell, i.e. translocating it through the cell membrane. This is immunologically highly advantageous, since the epitope will be brought more closely in contact with immunologically 25 competent cells of the body to which the fusion proteinexpressing vaccine cells are administered.

Secondly, the adhesin part of the epitope-carrying fusion protein can be selected so as to have specific binding properties whereby the vaccine cell may be targeted to a particu-30 lar location in the body where an immunological response to the epitope is desirable. The adhesion of the epitope-carrying cell to a particular location or region of the body will in this manner ensure that the cell is retained in the human

or animal body in that particular location for a period of time which is sufficient to obtain the desired immune response.

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In accordance with the invention, a useful cell for expression of the above fusion protein is one selected from a bacterial species which inherently contains an adhesin-carrying surface organelle. Such species include as examples gramnegative species of Enterobacteriaceae such as E. coli, Klebsiella spp, Salmonella spp, Yersinia spp, Vibrionaceae, Hemophilus spp, Bordetella spp and Pseudomonadaceae, and gram-positive species such as Neisseria spp and Streptococcus spp.

The epitope part of a fusion protein according to the invention can be an epitope derived from any pathogenic organism or agent against which it is desirable to develop vaccines. Such pathogenic organisms include viruses, bacteria and eucaryotic organisms such as fungi, yeast or protozoa.

Whereas cells expressing an epitope-carrying fusion protein as defined herein may be used as a live vaccine, it is also within the scope of the invention to provide isolated and/or purified cell surface organelles comprising the fusion protein, including fimbriae and pili, as a vaccine, and it is also contemplated that useful vaccines may be provided wherein cells expressing an epitope-carrying fusion protein have been killed by conventional methods such as formaldehyde treatment or thermal treatment.

In a further embodiment of the invention, the fusion protein according to the invention comprises as the non-adhesin polypeptide part a toxic gene product e.g. having a selective toxic effect on particular cells in the body such as e.g. cancer cells. By selecting the adhesin part as one having a specific binding affinity to receptors in such cells it is possible to have cells expressing the toxic gene product bound selectively to such target cells whereby these cells

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may be killed or damaged by the toxic gene product. It is also possible to use isolated or purified cell organelles containing a fusion protein comprising the cell toxic (cytotoxic) gene product for the purpose of targeting the toxic product.

In a further interesting embodiment, the fusion protein comprises an antibody. Such an embodiment is, inter alia, particularly interesting with respect to the provision of fusion proteins which may be used in affinity purification of biological compounds having binding affinity to the antibody part of the fusion protein. It is contemplated that cells expressing as part of a surface organelle, such a fusion protein may be utilized directly as a means of concentrating a biological compound, or the isolated surface organelles comprising the antibody-carrying fusion protein may be used for this purpose.

Furthermore, the fusion proteins as defined herein are useful as carriers of non-covalently bound compounds such as pharmacologically active, diagnostically active and imaging compounds with the purpose of providing cells or cell organelles carrying the active compounds, which thereby become targetable to particular regions or locations of a body to which these cells or cell organelles are administered. The invention encompasses any combination of a fusion protein as defined herein and an active compound which can be covalently bound to a fusion protein.

As mentioned above, the present invention encompasses in one aspect a recombinant bacterial cell which expresses a recombinant bacterial adhesin variant or a FimH adhesin as defined above. In one specific embodiment, the bacterial cell is one which comprises the above-defined recombinant replicon. Depending on the field of application of such a cell, it may e.g. be selected from a soil bacterium, an aquatic bacterium, a bacterium which is normally associated with plants, a bacterium which is member of the human or animal indigenous

bacterial flora, or a bacterium which is adapted to colonize certain ecological niches such as e.g. sewage purification plants or certain inanimate surfaces.

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The major significant advantages which have been achieved by the present invention is the possibility to provide recombinant bacterial cells which are not only ecologically welladapted to grow in a particular ecological environment, but which are also provided with means for colonizing more permanently in their ecologically natural environment. These means for improved ability to colonize an environment are the 10 adhesins expressed by the bacteria which have been constructed and/or selected so as to enable the recombinant bacterial cell to adhere to or bind to specific receptors in the environment, i.e. the bacterial cells are targeted to that environment. Thereby the bacteria according to the 15 present invention will have an ecologically competitive advantage relative to organisms in the particular environment which do not have surface structures comprising adhesins binding to receptors present in the environment, at least not to the same extent as the bacterial cells according to the 20 invention.

In addition to the environment-specific adhesins which the bacterial cell expresses, the cell will have a phenotype which is desirable in the environment to which it is targeted. As one example, a cell according to the invention 25 which is originally isolated from the human or animal indigenous bacterial flora may typically be one which expresses an epitope including an epitope which is part of a fusion protein expressed by the bacterial cell. As another example may be mentioned a bacterial cell which is isolated from a plant 30 and which expresses a pesticidally active compound such as a Bacillus thuringiensis toxin. Further examples include a plant root-associated nitrogen-fixating bacterium isolated from soil which in accordance with the invention is provided with adhesins improving the capability of the bacterium to 35 become permanently colonized to the roots of a specific plant or specific plants, or a bacterium which is ecologically associated with an aquatic or terrestrial environment containing pollutants to be degraded or removed.

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Accordingly, the recombinant bacterial cell can be derived from any gram-negative or gram-positive bacterium for which a need exists to obtain improved colonization in a particular inanimate or animate environment. Such bacteria include as examples Enterobacteriaceae spp, Hemophilus spp, Neisseria spp, Bordetella spp, Streptococcus spp, Pseudomonadaceae spp, Vibrionaceae spp, Baccilaceae spp.

In certain embodiments of the invention it is advantageous that the present recombinant bacterial cell is provided as one which, when it is administered to a particular location or environment, will not persist in that environment. Accordingly, such a recombinant bacterial cell may further comprise 15 a gene coding for a gene product which, when expressed has a killing or cell function-limiting effect in said cell, the $^{\circ}$ expression of said gene coding for the cell killing or cell function-limiting gene product being regulated in such a manner that the bacterial cell when targeted to receptor in a 20 specific location will be killed or limited in its function in a pre-determined manner. The gene coding for the cell killing or cell function-limiting gene product is suitably regulated by a factor selected from the group consisting of a stochastic event, the presence/absence of a chemical compound 25 in the location, and a physical factor.

In a further aspect, the invention relates to a method of isolating or constructing a recombinant bacterial cell expressing an adhesin having modified binding properties relative to a natively expressed adhesin such as a natively expressed FimH adhesin. As it is defined above, this method comprises identifying in the bacterial cell DNA sequence(s) coding for the binding domain(s) of said natively expressed adhesin and substituting at least one codon herein whereby a modified adhesin molecule is expressed that is different in

at least one amino acid from the adhesin expressed natively, and selecting a bacterial cell expressing the modified adhesin having an altered adhesion phenotype relative to the natively expressed bacterial adhesin.

As it is explained in details below, the binding domain can e.g. be identified by constructing chimeric adhesin-encoding genes and screening for cells which by having a region in the adhesin gene replaced by a corresponding heterologous region of a different DNA sequence, acquires a new binding phenotype. Having identified a binding domain of the natively 10 expressed adhesin, recombinant cells having desirable binding phenotypes may be obtained by substituting one or more codons in the binding domain(s) to obtain expression of recombinant adhesins and selecting cells having the desirable phenotypes. The substitution of codons may be achieved by methods know 15 per se such as site-directed mutagenesis using synthetic oligonucleotides and PCR technology or transposable elements or by conventional radiation or chemical mutagenization.

In certain useful embodiments, the above method includes

steps whereby a non-adhesin compound is associated with the adhesin, e.g. a step where a gene coding for the recombinant adhesin is part of a hybrid gene comprising a gene coding for a non-adhesin polypeptide which thereby is expressed with the recombinant adhesin as part of a fusion protein comprising

the adhesin. Furthermore, recombinant adhesins resulting from the above method may in specific embodiments comprise a non-covalently bound compound which is associated with the adhesin when expressed.

As mentioned above, the invention also encompasses recombinant bacterial cells having selected binding properties
whereby cells with desirable phenotypes can colonize environments where the presence of bacteria having a particular
phenotype is advantageous. Accordingly, there is in a further
aspect of the invention provided a method of preparing a
recombinant bacterial cell that binds to a specific receptor

moiety, comprising introducing into a bacterium that does not produce an adhesin binding to said receptor moiety, a DNA sequence coding for an adhesin binding to the receptor moiety, and selecting a bacterial cell expressing the DNA sequence.

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The primary objective of this method is to provide the means of constructing a bacterial strain having the capacity to colonize an environment, based on a parent strain which has an inherent, useful phenotype in this particular environment but which does not express an adhesin binding to receptor 10 moieties in the environment. Accordingly, the method includes as a first step the isolation of an environmentally adapted bacterium not binding to appropriate receptor moieties and in subsequent steps, the identification of heterologous genes encoding adhesins which bind to receptor moieties occurring 15 in said environment, preferably moieties occurring abundantly, isolating this gene and introducing it into the above parent strain. The adhesin gene may e.g. be a gene coding for a naturally occurring FimH adhesin or a recombinant FimH adhesin as defined above. 20

In one useful embodiment of the method, the adhesin-encoding gene is introduced by transforming a parent bacterial cell with a recombinant replicon as defined herein. In further embodiments, the method is designed so as to obtain a cell wherein a non-adhesin compound is associated with the adhesin, e.g. by introducing the gene coding for an adhesin as a hybrid gene coding for a non-adhesin polypeptide whereby non-adhesin compound is expressed with the adhesin as part of a fusion protein comprising the adhesin, or by binding non-covalently a compound to the adhesin when expressed.

Besides the above method, an adhesin carrying bacterial cell having an altered pattern of adhesion can be provided by using a selection procedure comprising contacting an appropriately sized population of wild-type adhesin-carrying bacterial cells with a potential receptor moiety to which the

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wild-type cells do not adhere, e.g. in a manner as it is disclosed in Example 6 below whereby spontaneously or randomly mutated cells having acquired the ability to adhere to the receptor moiety in question, become progressively enriched. From such an enriched culture, cells with the new adhesion ability can readily be isolated and further characterized.

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As it has been explained in details above, one primary objective of the present invention is to provide the means of targeting a compound to a specific location. Accordingly, the invention relates in an important aspect to a method of targeting an adhesin to such a location. The method comprises the identification in the location of a receptor moiety, said moiety preferably being one which occurs abundantly in the particular location, which moiety can recognize and interact with an adhesin, and the isolation of a bacterial cell which is capable of growing in the location and expressing an adhesin which recognizes and interacts with the identified receptor moiety, and administering the cell or the adhesin in an isolated form to that location.

The identification of a suitable receptor moiety in a particular location can be carried out in several manners. One example is a screening procedure where cells expressing known adhesins or known isolated adhesins are administered to the location e.g. being isolated cells or tissues of microbial, animal or plant origin or an inanimate surface as defined herein, and screening for binding/adhesion of the tested adhesins e.g. according to adhesion assays as disclosed herein. If binding of one or more adhesins occurs, it is an indication that receptor moieties for that or those tested adhesin(s), is/are present in the location.

Alternatively, available data with regard to the presence and amounts of chemical moieties present on the surfaces of the location may be collected or such data have to be generated, and based upon such data, adhesins which are known to bind to one or more of the identified major moieties are selected and

their binding to this/these structure(s) is tested e.g. according to the assays as used herein. Chemical moieties which are considered potential adhesin-interacting receptor moieties include as examples glycolipids, glycoproteins, proteins, polypeptides, saccharide moieties and peptides.

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In the case no suitable chemical moiety is identified in the location, which is capable of binding to known adhesins or which bind with a sufficient affinity, it is required to construct a library of modified adhesin molecules based on known adhesins which are modified by replacing one or more 10 codons as it is explained herein, and/or such a library provided by constructing synthetic adhesin molecules, and then screening this library for recognition of and interaction with identified location surface moieties. A library of modified FimH adhesins may e.g. be selected for specificity towards a given receptor by running clones of these adhesins through a column or matrix containing the receptor moiety in question or cells or tissues isolated from the location without knowing what the receptor moiety is. The clone(s) expressing the adhesins with affinity to receptor moie-20 ty/moieties will adhere/bind to the column or matrix, and can subsequently be isolated therefrom.

It is within the contemplation of the invention that crystallographic analyses of adhesins, whether naturally occurring or constructed as indicated above, is a useful technique for the obtainment of information about adhesin structures that assumingly will recognize and interact with particular adhesin receptor moieties.

In accordance with the invention, one embodiment of the above method is one wherein the isolated bacterial cell expresses an adhesin having modified receptor moiety-binding properties relative to an adhesin natively expressed by the cell, the isolation of the cell comprising identifying in a parent bacterial cell, DNA sequence(s) coding for the binding domain(s) of said natively expressed adhesin and substituting

at least one codon herein, whereby a modified adhesin molecule is expressed that is different in at least one amino acid from the adhesin expressed natively, and selecting a bacterial cell expressing the modified adhesin having an altered adhesion phenotype relative to the natively expressed bacterial adhesin or a method wherein the bacterial cell expressing an adhesin that recognizes and binds to the receptor moiety is a recombinant bacterial cell derived from a parent bacterial cell that does not produce an adhesin binding to said receptor, by inserting into the parent cell a DNA sequence coding for an adhesin binding to the receptor moiety, and selecting a bacterial cell expressing the DNA sequence.

One primary objective of the present invention is the targeting of useful non-adhesin compounds to a particular location.
Accordingly, the invention encompasses in an interesting
embodiment a method as defined above wherein a non-adhesin
compound is associated with the adhesin, whereby said compound is targeted with the adhesin to the location comprising
the receptor moieties recognizable by the adhesin.
The compound can be associated with the adhesin by a covalent
binding or by any of the above mentioned non-covalent types
of molecule interaction forces.

When associated covalently with the adhesin the compound to

be co-targeted to the selected location with the adhesin can
be an enzyme, an antibody, an epitope or a toxin which is
part of a fusion protein comprising the adhesin. A compound
which is associated with the adhesin by a non-covalent binding is typically a pharmacologically active, diagnostically
active or imaging compound.

Locations to which it is desirable to have an adhesin targeted by the present method include a human or animal surface, a plant surface and an inanimate surface as defined above.

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In one specific embodiment of the present method the bacterial cell being administered to the location expresses a recombinant bacterial adhesin variant derived from a naturally occurring parent adhesin, said adhesin variant having altered binding properties relative to the naturally occurring adhesin from which it is derived, the altered binding properties including binding to at least one receptor moiety to which the parent adhesin does not bind. Such an adhesin variant is advantageously derived from a naturally occurring adhesin isolated from a cell structure selected from the group consisting of a capsule, a lipopolysaccharide layer, on outer membrane protein, a flagellum, a pilus, a fimbria, a non-fimbrial adhesin (NFA) or an afimbrial adhesin (AFA).

In specific embodiments of the invention, the above adhesin variant as used in the present method is a protein having an 15 amino acid sequence differing in at least one amino acid residue from its parent protein adhesin such as a FimH adhesin having an amino acid sequence which differs from the E. coli PC31 FimH adhesin as defined herein in at least one amino acid. Such a FimH adhesin includes an adhesin which 20 binds to a receptor selected from the group consisting of a domain where mannosyl residues are not terminal and a domain devoid of saccharide and an adhesin variant which is at least 90% homologous to the PC31 FimH adhesin as defined herein, such as at least 92% homologous, more preferably at least 93% 25 homologous, even more preferably at least 94% homologous, most preferably at least 95% homologous, and in particular at least 96% homologous, e.g. at least 97% homologous. In particularly interesting embodiments, the adhesin is at least 98% homologous, including at least 99% homologous or at least 30 99.5% homologous.

The above FimH adhesin can be a chimeric adhesin as defined above, comprising amino acid sequences from different FimH adhesins and constructed according to the methods below.

In accordance with the invention, an adhesin can be administered to a location in the form of an adhesin-expressing bacterial cell. Such a cell is one capable of growing in that particular location. Accordingly, the bacterial cell is suitably derived from a bacterial species which is normally occurring in the location including human or animal body surfaces, plant surfaces such as plant root surfaces and inanimate surfaces. In this context, an animal body surface includes the insect gut, whereto it is desirable to administer a bacterial cell expressing an insecticidally active toxin.

Thus, if it is desired to administer the bacterial cell to the root of a plant, a suitable bacterial cell is preferably isolated from a strain which has colonized the rhizosphere of that plant to a large degree, i.e. the strain is a major member of the natural plant root flora. Such an isolate is then provided with a gene coding for an adhesin which will recognize and interact with an abundantly occurring moiety on the roots of said plant. In this manner, a suitable adhesin which is expressed naturally in a bacterium which is not adapted to grow in a plant rhizosphere, becomes expressible in a normal inhabitant of the rhizosphere environment (location).

In specific embodiments of the present method of targeting a bacterial adhesin to a specific location, the adhesin is a FimH adhesin as defined above, having an amino acid sequence which differs from the *E. coli* PC31 FimH adhesin as defined herein in at least one amino acid.

In an interesting embodiment, the adhesin-carrying bacterial cell being targeted is a cell which further comprises a gene coding for a gene product which, when it is expressed, has a killing or cell function-limiting effect in said cell, the expression of said gene coding for the cell killing or cell function-limiting gene product being regulated in such a manner that the bacterial cell, when targeted, will be killed

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or limited in its function in a pre-determined manner. The expression of such a "suicide" or cell function-limiting gene may suitably be regulated by a factor selected from the group consisting of a stochastic event, the presence/absence of a chemical compound in the location and a physical factor. As examples of such "suicide" or cell function-limiting genes providing the means of biological containment, may be mentioned those disclosed in WO 87/5932 and WO 93/20211

Furthermore, the present Designer Adhesin Technology (DAT) provides very useful means of obtaining colonization with 10 desirable bacteria in a particular environment with the purpose of obtaining beneficial changes of the microbial flora in the environment. As one example, certain bacterial species in the gastrointestinal (GI) tract of humans and animals have beneficial effects on the health condition of 15 the host organism e.g. by suppressing pathogenic organisms or by contributing to the digesting of certain diet components. The present technology makes it possible to select particularly useful bacteria from the GI-tract and have them 20 designed in accordance with the present invention, to have improved colonization abilities. Similar examples include desirable bacterial colonizations of biological sewage purification systems, plants where invasion of pathogenic organisms may be controlled by colonizing the plants with harmless bacteria, and teeth where caries may be controlled by coloni-25 zing the dental enamel with bacteria suppressing those causing the caries attacks.

In another industrially interesting aspect, the invention provides the means of isolating a compound from a solution or suspension containing the compound. The method comprising contacting the solution or the suspension with a fusion protein as defined herein wherein the heterologous polypeptide has an affinity to the compound to be isolated.

Furthermore, the invention provides a composition comprising a population of a bacterial cell as defined herein.

The invention is further illustrated in the below Examples and the Figures, wherein

Fig. 1 is a schematic model for the construction of recombinant plasmids pGBl-24 (containing fimH from CI #10) and pGB2-24 (containing fimH from PC31) used for transforming E. coli AAEC191A(pPKLll4) with cloned fimH genes. Plasmid pGB2-24 was used as the vector for all other cloned fimH genes described herein;

Fig. 2 is a restriction map of fimH genes. Five unique restriction sites are present in the PC31 fimH gene. Numbers in parentheses following enzymes are the base pair positions of the cut sites. Some of these sites are found in the other fimH genes, as marked. Chimeric genes were produced by exchanging each available restriction fragment from the other five fimH genes with corresponding fragments in the PC31 gene and then recombinant strains expressing resulting chimeric fimH subunits were tested for adhesion. Fragments indicated by boxes are those which conferred MF or MFP adhesive phenotypes on the chimeric genes;

Fig. 3 illustrates adhesion of representative "wild-type" (A) 20 and recombinant (B) M-class, MF-class and MFP-class strains to Mn (1), Fn (2), periodate-treated Fn (3) and to FnSp1 (4). Strain designations given for the "wild-type" strains are given in AS. Strain designations KB31, KB12, KB4, KB7, KB50 and KB10, are for recombinant strains of AAEC191A(pPKL114), 25 which is fimH, after transformation with plasmids that contain fimH+ from strains HB101(pPKL4), CI #12, CI #4, CI #7, CSH-50 and CI #10, respectively. Open columns indicate results when bacteria were incubated in buffer without Dmannose, while solid columns are results in the presence of 30 D-mannose. Values indicated are the mean \pm S.E.M. (n=4) for each column;

Fig. 4 illustrates the adhesion of representative M-class, MF-class and MFP-class strains (CIs #12, #4 and #10, respect-

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ively) to Fn fragments prepared by thermolysin treatment as described in ref. 51. Columns labelled 1-5 indicate adhesion to: 1) NH2-terminal 30-kDa domain; 2) the 55-kDa gelatin-binding domain; 3) the 110-kDa cell attachment domain; 4) the 29-38-kDa heparin binding domains; and 5) the 20-kDa COOH-terminal domain. Open columns represent adhesion in the absence of D-mannose; solid columns represent adhesion in the presence of D-mannose. Mean ± S.E.M. (n=4);

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Fig. 5 is a composite figure illustrating comparison of amino acid sequences of FimH adhesins and active restriction frag-10 ments of fimH genes. The published nucleotide and deduced amino acid sequence of the PC31 fimH gene and gene product (ref. 27) serve as prototype. Numbered amino acid residues shown above the model of the PC31 FimH represent residues that are different in other FimH subunits due to amino acid 15 substitution or deletion. Standard one-letter code applies and residues in the other FimH sequences that are different are indicated. Deleted amino acids are indicated by Δ . It should be noted that residue 176 is not arginine as published previously (ref. 27) for the PC31 FimH, but proline. Regions 20 of the FimH subunits conferring change in adhesive phenotype, highlighted in bold, were determined by functional assays performed on chimeras between the "classic" mannose-specific PC31 fimH gene present in HB101(pPKL4) and the above described genes. Residues predicted to be key in conferring 25 receptor specificity are circled. Approximate positions of unique restriction sites used to create chimeras are indicated along the bottom of the model;

Fig. 6 illustrates plasmid pPKL4 which is a derivative of pBR322 (thick line) carrying the entire fim operon (FimA-H) including the regulatory genes fimB and fimE (not shown), and the promoter region with the SnaBI site. In this plasmid an 8mer linker with an BglII site was inserted in the SnaBI site to create pPKL83;

Fig. 7 illustrates the construction of plasmid pSM1314; the vector pVLT33 is a derivative of the broad host range replicon RSF1010. Plasmid pPKL83 was digested with BglII and pVLT was digested with BamH1; the two were ligated and pSM1314 was the resulting plasmid in which expression of the fimA-H cluster is under the control of the tac promoter;

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Fig. 8 illustrates plasmid pLPA22 and derivatives hereof as used in this study. The triangles indicate the position of translational stop-linkers in the fimH gene in plasmid pPKL115. The positions of heterologous inserts are indicated (black boxes). Small triangles indicate signal-peptide encoding sectors.

Fig. 9 illustrates plasmids pLPA29, pLPA30, pLPA36, pLPA58, pLPA59 and pLPA98;

Fig. 10 shows immuno-electron microscopy with colloid gold labelling of $E.\ coli$ HB101 cells containing plasmids pLPA22 plus pPKL115 (a), pLPA37 plus pPKL115 (b), pLPA38 plus pPKL115 (c), using anti-pre-S2 monoclonal antiserum. Bar, 0.1 μ m.

20 EXAMPLE 1

Functional heterogeneity of type 1 fimbrial adhesins due to minor sequence variations among fimH genes

1.1. Materials and methods

1.1.2. Reagents

Yeast Mn, a polymannosylated glycoprotein isolated from Saccharomyces cerevisiae cell walls, was obtained from a commercial source (Sigma Chemical Co, St. Louis, MO, U.S.A.). Mannan is composed of an N-linked backbone of β 1,2-linked mannopyranose units with α -linked mannopyranose side chains (ref. 38). The majority of the carbohydrate of human plasma

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Fn is composed of N-glycosidic complex-type biantennary glycans and no high mannose-type or hybrid-type N-glycans have been described (refs. 30, 45, 54). Human plasma Fn and Fn fragments were purified as described previously (refs. 5, 15, 51, 58). Periodate treatment was performed as described previously (ref. 51). The synthetic peptide, FnSpl, copying the first 30 amino acid residues of the Fn molecule (EAQQMVQ-PQSPVAVSQSKPGCYDNGKHYQI) was synthesized in the Protein Chemistry Laboratory of the VA Medical Center, Memphis, TN (SEQ ID NO:2). The saccharide content of the four substrata was characterized using two lectins, concanavalin A (ConA), well known to react with terminal and internal mannosyl residues, and the Calanthus nivalis agglutinin (GNA), which recognizes only terminal Manlphal-3Man, Manlphal-6Man and Manlphal-2Man sequences (E. Y. Laboratories, San Mateo, CA). Immobi-15 lized Mn and Fn both reacted with ConA, whereas GNA bound only to Mn. These results are consistent with the known structures of the oligosaccharide moieties of these two compounds. Neither lectin reacted with immobilized FnSpl. Periodate treatment (ref. 51) of Mn or Fn eliminated lectin 20 reactivity.

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1.1.3. Bacterial strains and plasmids

The CSH-50 strain (lambda, FaraΔ(lac-pro) rspL thi fimE::IS1) is a Cold Spring Harbor K12-derived strain (ref. 35). The E. coli strain MG 1655 (CGSC6300; K12 derivative, lambda, F) and a derivative strain AAEC191A (MG1655 recA Δfim were generously provided by Dr. Ian Blomfield (Bowman Gray University, Winston-Salem, NC). AAEC191A has had the entire fim gene cluster deleted by allelic exchange (ref. 8).

Clinical isolates (CIs) were urinary tract isolates obtained from the clinical microbiology laboratories of the Memphis VA Medical Center or The City of Memphis Hospitals, Memphis, TN. The 12 CIs used in this study were selected on the basis of MS agglutination of yeast cells after growth in broth, a classic test for type 1 fimbriae.

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Plasmid pPKL4, a pBR322 derivative containing the entire fim gene cluster from $E.\ coli$ strain PC31 (K12-derivative, gal tonA phx ara) and encoding for the expression of fully functional type 1 fimbriae in HB101 (supE hsdS recA ara proA lacY galK rspL xyl mtl Δ fimBE), has been described previously (ref. 28). pPKL114 is a recombinant plasmid derived from pPKL4, but with a translational stop-linker inserted into the Kpnl site in the fimH gene. No transcriptional effects of the

stop-linker are to be expected. Antibiotics were used at the following final concentrations: ampicillin (50 μ g/ml), kanamycin (60 μ g/ml) and chloramphenicol (30 μ g/ml).

1.1.4. Polymerase chain reaction

Oligonucleotide primers were designed using the published sequence for the fimH gene in pPKL4 (ref. 27). The 5' primers 15 copied regions 13 and 49 bp upstream from the fimH gene and were extended on the 5' end by an Apall restriction site and a GC clamp: Primer 1: 5'-GGGGG-GTGCAC-ACC TAC AGC TGA ACC CGG-3' (SEQ ID NO:3); Primer 2: 5'-GGGG GTGCAC T CAG GGA ACC ATT CAG GCA-3' (SEQ ID NO:4). The 3' primers copied 18 bases 20 of the bottom strand of the fimH gene that encode for the 6 terminal amino acids of fimH and were extended by an Fspl or Sph1 site and a GC clamp: Primer 3: 5'-GGG TGCGCA TTA TTG ATA AAC AAA AGT CAC - 3' (SEQ ID NO:5); Primer 4: 5'-GGG GCATGC TTA TTG ATA AAC AAA AGT CAC-3' (SEQ ID NO:6). Primer 1 and 3 were used for CI #10 and pPKL4, primer 1 and 4 were 25 used for CI #4 and CSH-50 and primer 2 and 4 were used for CI #s 7 and 12 to generate PCR products from plasmid or chromosomal DNA prepared from E. coli expressing different functional classes of type 1 fimbriae. The PCR reaction 30 mixture consisted of template DNA, primer pairs, dNTPs, and Taq DNA polymerase in PCR buffer. The PCR was performed in a Perkin-Elmer Cetus automatic thermal cycler with denaturation at 96°C for 1 min., primer annealing at 55°C for 1 min., and primer extension at 72°C for 2 mins. for a total of 40 cycles. All of the PCR products migrated similarly in agarose 35

gels. Purification, restriction and ligation of DNA was performed using standard procedures (refs. 39, 48). All primers for PCR and for nucleotide sequencing were produced by the Molecular Resources Center, UT, Memphis.

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5 1.1.5. Subcloning

The PCR products from CI#10 and from pPKL4 were cut with respective restriction enzymes and ligated into the Apall and Fspl restriction sites of plasmid pACYC177 (New England Biolabs, Beverly, MA, U.S.A.) which is compatible with the pBR322-based pPKLll4 to be used in complementation experi-10 ments, creating plasmids pGBl and pGB2, respectively (Fig. 1). However, it became inconvenient to use pACYCl77-based plasmids because of a high frequency of appearance of spontaneous Kmr in the AAEC191A host strain. The origin of this 15 problem is not entirely clear, but it was avoided by subcloning the fimH genes from pGBl and pGB2. The inserts and upstream regions of pACYCl77 containing the tet promoter were cut from pGB1 and pGB2 with Fspl and BamH1 and subcloned into the polylinker site of pGEM-3Z (Promega, Madison, WI) that 20 had been cut with BamHl and Hinc2, creating plasmids pGBl1 and pGB2-1 respectively. pGEM-3Z was simply used as a convenient intermediate in subcloning into pACYC184.

The inserts were cut out again using Smal and Hind3 and subcloned into pACYC184 (New England Biolabs, Beverly, MA) cut with Hinc2 and Hind3, creating plasmids pGB1-2 and pGB2-25 24 containing the fimH genes from CI#10 and pPKL4, respectively. These plasmids complement the non-adhesive defect of AAEC191A(pPKL114) giving the adhesive phenotypes of the parental strains (see Results). Cutting the fimH gene from pGB2-24 using Apall and Sph1 makes it possible to easily insert other fimH genes obtained by amplifying chromosomal DNA of other isolates by PCR. All recombinant strains we have tested thus far using this technique exhibit the same adhesive phenotype as the parent strains from which the fimH genes were cloned.

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1.1.6. Construction of chimeric fimH genes

Unique restriction sites (Fig. 2) were used to construct chimeric fimH genes between the prototypical MS pPKL4 fimH gene, used as genetic background, and restriction fragments obtained from the newly described fimH genes. Fragments were purified from agarose gels and ligated into restriction "spaces" generated in the pPKL4 fimH gene present in pACYCl'84 (pGB2-24). Each chimera was analyzed by restriction mapping and the nucleotide sequences of bridging segments were determined to ensure proper constructions. The plasmids containing chimeric fimH genes were transformed into AAECl91A(pPKL114) and clones were tested for agglutination of yeast cells and for adhesion to Mn, Fn and FnSp1.

1.1.7. Nucleotide sequencing

The nucleotide sequences of fimH genes were determined by the 15 dideoxynucleotide chain termination method of Sanger (ref. 49) using a Sequenase II® kit (U.S. Biochemical Corp., Cleveland, Ohio) and $[\alpha^{-35}S]dATP$ (800 to 1,000 Ci/mmol) according to the manufacturer's suggestions. The amino acid sequences were deduced from nucleotide sequences using MacVector® 20 protein and DNA analysis software (Eastman Kodak, Rochester, NY). To ensure fidelity of the PCR amplification, selected fimH genes were re-amplified, cloned, tested for activity and re-sequenced. More recently, we have used the $fmol^m$ Polymerase Sequencing System (Promega, Madison, WI), because it is 25 useful with small amounts of DNA and thus subcloning the fimH genes from the pACYCl84-based plasmids to high copy number plasmids was obviated. Bands were visualized by autoradiography of sequencing gels and compared with the published fimH gene sequence (ref. 27). 30

1.1.8. Yeast cell aggregation assay

E. coli were tested for their ability to aggregate yeast cells. Commercial baker's yeast, Saccharomyyces cerevisiae,

was suspended in PBS (5 mg dry weight/ml). E. coli were washed in PBS, resuspended to an OD₅₃₀ of 0.4, and mixed with the yeast cell suspension in PBS with or without 1% D-mannose. Aggregation was monitored visually and the titer recorded as the last dilution giving a positive aggregation reaction.

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1.1.9. Adhesion assays

Adhesion assays were performed as described previously (ref. 51). Briefly, microtiter assay wells were coated with

10 receptor molecules as indicated in the text and figure legends. After the wells were washed two times with PBS, 100 µl bacterial suspensions were added in 0.1% BSA-PBS. After incubation at 37°C for indicated times, wells were washed three times with PBS and adherent bacteria were detected by using rabbit anti-E. coli serum. Antibody binding was detected using peroxidase-conjugated goat anti-Rabbit IgG. Reaction product generated from the 5-aminosalicylic acid substrate was measured at 405 nm after 10-15 minutes by using an automatic microplate reader (Molecular Devices, Inc.,

20 Menlo Park, CA). Values reported are corrected for background reaction using BSA coated plates as control.

1.2. Results

In a previous publication it was reported that type 1 fimbriae of E. coli CSH-50 and HB101(pPKL4) differ func
25 tionally in their pattern of adhesion to Mn, Fn, periodatetreated Fn and a synthetic peptide, FnSp1, immobilized on plastic (ref. 51). Since CSH-50 and HB101(pPKL4) are laboratory strains, we tested 12 clinical E. coli isolates (CIs) obtained from human urine for adhesion to these four substra
30 ta. All of the CIs agglutinated yeast cells in a MS fashion. Five of the twelve CIs adhered only to Mn. The adhesive activity of HB101(pPKL4) and of CI #12 are shown as examples of this class, which we have tentatively designated as M class (Fig. 3A). Three of the 12 CIs adhered to Mn and Fn,

but not to periodate-treated Fn or to FnSp1. The adhesive activities of CI #s 4 and 7 are shown as examples of this class, designated as MF class. Three of the twelve CIs adhered to each of the substrata. The adhesive activities of CSH-50 and CI #10 are shown as examples of this class, designated as MEP class.

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Adhesion of strains representing these three classes to Fn fragments further illustrates the distinct differences between the three classes. The M class CI #12 does not adhere to any of the Fn fragments (Fig. 4). The MF class CI #4 adheres to the 40-kDa gelatin-binding fragment. The MFP class CI #10 adheres, with only slight differences, to all 5 fragments of Fn tested. Periodate treatment eliminated binding of CI #4 to domain 2, but had no effect on the binding of CI #10 to any of the Fn domains (data not shown).

Since the fimH subunit has been shown to mediate the mannose-sensitive activity of type 1 fimbriae, we focused our initial efforts to elucidate the molecular basis for the observed functional heterogeneity on the fimH gene. fimH genes were amplified from chromosomal (or plasmid, for pPKL4) DNA and the genes were cloned into pACYC177 and subcloned into pACYC184 under control of the β -lactamase promoter of pACYC177, according to Materials and Methods (Fig. 1)

The adhesive phenotypes conferred by the fimH genes were

25 tested in the following way. E. coli K-12 strain AAEC191A
(Δfim) was first transformed with plasmid pPKL114, which
contains an intact fim gene cluster but with a translational
stop-linker inserted into the last gene, fimH. This derivative produces morphologically normal fimbriae that are non30 adhesive due to absence of the FimH subunit. Plasmids harbouring cloned fimH genes were transformed into E. coli
AAEC191A(pPKL114) and the resultant strains were tested for
their ability to adhere to Mn, Fn, periodate-treated Fn and
to FnSpl (Fig. 3B). Each of the recombinant strains displayed
35 adhesive phenotypes mimicking those of the representative

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parent strains from which the fimH genes were obtained. fimH genes were cloned from each of the other 8 CIs and similar results were obtained with the adhesion of recombinant strains mimicking that exhibited by the parental CIs.

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The complete nucleotide sequences of each of the six representative fimH genes were determined and the amino acid sequences of the fimH proteins were deduced as it is shown in Table 1 below which is a representation of amino acid sequences of the FimH subunits deduced from nucleotide 10 sequences of selected fimH genes disclosed in this example [CI#3 (SEQ ID NO:33), CI#4 (SEQ ID NO:29), CI#7 (SEQ ID NO:30), CI#10 (SEQ ID NO:31) and CI#12 (SEQ ID NO:28)] and those of the E. coli K12 strain PC31 (SEQ ID NO:1) and E. coli strain CSH-50 (SEQ ID NO:32). Additionally, the FimH amino acid sequences of the following clinical isolates of E. coli are shown: KB21 (SEQ ID NO:27), KS54 (SEQ ID NO:35), U221-3 (SEQ ID NO:36), MJ#9-3 (SEQ ID NO:37), MJ#31-3 (SEQ ID NO:38), MJ#11-2 (SEQ ID NO:39), MJ#2-2 (SEQ ID NO:1) and F-18 (SEQ ID NO:34). Standard one-letter code applies. Deleted amino acid residues are indicated by $\Delta s.$ M, M^L , MF, MFP, and 20

 ${ t M}^{ t R}$ indicate the adhesin class as defined above.

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Table 1. Amino acid sequences of the FimH proteins deduced from nucleotide sequences of fimH genes of clinical isolates disclosed in this example and of E. coli K12 strains PC31 and CSH-50

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and gene products should also be identical, subunit incorporation into the fimbrial superstructure should not vary significantly. These results emphasize that in these experiments it is the FimH subunit that determines receptor specificity.

In comparing the new FimH sequences to the one published previously (ref. 27), the only structural alteration that can be clearly linked to a functional change, without resorting to analysis of chimeric fimH genes, is the non-conservative substitution of arginine⁵⁸ in the MFP class CSH-50 FimH subunit for leucine⁵⁸ in the M class PC31 FimH subunit. Since each of the other FimH sequences had more than one change, it was necessary to construct chimeric genes to begin to focus on functionally relevant changes.

In the case of the CI #10 FimH, an MFP class adhesive activity similar to that of CSH-50 is conferred by a different region of the gene which encodes for a subunit deleted of residues 116-119. It remains to be determined how two distinctly different structural changes can bring about apparently similar changes in receptor specificity. It is possible, of course, that as additional receptor molecules are tested, these two variants will be found to be functionally distinct.

The Apal1-Tth111I fragment of the CI #7 fimH gene confers MF class activity in the CI#7/PC31 fimH chimera. Since the asparagine 16-threonine 16 substitution is within the leader 25 sequence and thus not represented in the mature protein, the histidine³³- asparagine³³ substitution must be of functional importance for the MF class CI #7 FimH. Comparison of the active regions of the MF class CI #4 and the M class CI #12 FimH subunits suggests the importance of the glutamic acid^{73} -30 ${\tt glycine}^{73}$ substitution for MF class activity of the CI #4 FimH. Thus, histidine 33, arginine 58, glutamic acid 37 and deleted glycine 116-isoleucine 119 appear to be key residues in the functional activity of FimH subunits of CI #7, CSH-50, CI #4 and CI #10, respectively, but a more precise demonstration 35

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of which residues are involved and how they affect the ligand-binding cleft(s) remains to be performed.

At first glance, the FimH mediated, mannose-sensitive protein-binding activity of type 1 fimbriae is the most surprising of the adhesive phenotypes described here. However, protein-binding activity of FimH (i.e. PilE) subunits was noted earlier in a study characterizing mutT-induced mutations in the fimH (pilE) gene (Harris et al., ref. 22) However, the protein-binding activity described by Harris et al. was not mannose-sensitive. It is presently not known whether 10 the protein-binding activity described herein is in addition to or separate from the mannose-binding activity, but the concept of bifunctional properties of lectins has been established for several years (ref. 6). While the MFP class type 1 15 fimbriae appears to react somewhat promiscuously with most Fn fragments, the reaction does not appear to be non-specific. For instance, the MFP class CSH-50 type 1 fimbriae do not adhere well to gelatin (ref. 51). Further, the adhesion to ovalbumin is sensitive to both periodate and glycosidase 20 treatment (ref. 51). Further work will be required to determine the consensus amino acid motif reactive with this class of FimH subunit.

Previous studies suggested that the combining site of the E. coli FimH adhesin is in the form of an extended pocket cor-25 responding to the size of a trisaccharide with an associated hydrophobic region (ref. 16). The MS nature of all of the adhesive interactions described suggests that if the combining sites are separate, they may be close to each other. However, it remains to be determined whether or not the mannose effect is direct or allosteric. Conformational changes that frequently occur in lectins upon binding the saccharide ligand (ref. 46) could affect a second, distant binding site. Site-directed mutations may be sufficient to clarify which structural changes result in changes in 35 receptor specificity. However, such studies are unlikely to shed much light on how the structural changes actually relate

Table 1, continued

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The nucleotide and deduced amino acid sequences of the pPKL4 fimH gene are identical to that reported previously, except that residue 176 is not an alanine residue as previously reported, but a proline residue. Independent re-amplification, re-cloning and re-sequencing confirmed this sequence for the pPKL4 fimH gene. Sequencing was also repeated on independently amplified and cloned isolates of the CI #10 and CI #7 fimH genes to confirm sequence fidelity and no errors

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The nucleotide and deduced amino acid sequences of the other 10 fimH alleles described in this Example are > 98% conserved, but there is more than one amino acid residue difference in all but one of the new fimH sequences when compared to the published pPKL4 sequence. To focus on the sequence differences that resulted in changes in functional activity, advan-15 tage of unique restriction sites were taken (Fig. 2) to construct chimeric fimH genes. Multiple restriction fragments covering the entirety of each of the sequenced fimH genes were exchanged with corresponding fragments in the prototypical fimH gene of E. coli strain PC31 that was amplified from 20 pPKL4, cloned into pACYCl84 and used as the genetic background. Recombinant plasmids containing the chimeric fimH genes were transformed into E. coli AAEC191A(pPKL114) and transformants were tested for adhesive phenotype, allowing determination of the regions of each gene capable of confer-25 ring functional activity (Fig. 5). All of the sequence changes that affected function occurred between residues 33

1.3. Discussion

were found.

The functional heterogeneity which is described above must be due entirely to allelic variants of the fimH gene. The only variables in the recombinant strains which are described in this Example are the fimH genes; all other genes necessary for fimbrial subunit synthesis, transport and assembly are the same in each case. Since the ratios of the various genes

and 119 of the 279 residue mature fimH sequence.

to the ligand-binding cleft(s) and it will ultimately be necessary to determine the 3-dimensional structure of FimH or FimH fragments crystallized in the presence of ligand to fully understand structure/function relationships.

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The three adhesive classes of type 1 fimbriae identified above may understate the functional heterogeneity of type 1 fimbriae. The group of CIs that has been tested in this Example is small and only a few substances have been tested as potential receptors. A larger group of isolates tested against additional receptor candidates might yield additional functional classes. Preliminary studies with MS Enterobacter aerogenes and Klebsiella pneumoniae strains exhibiting MF class and MFP class activity suggest that heterogeneous receptor specificities will also be found among other type 1 fimbriated enterobacterial species.

It is also believed that it is possible that adhesins from some fimbriae responsible for mannose-resistant hemagglutination or adhesion are structurally related to FimH, but with sequence alterations that eliminate sensitivity to mannose. The possibility that the MS lectin-like properties of FimH might be eliminated while retaining other adhesive properties of FimH (e.g. pellicle formation) has been shown previously in a study characterizing mutT-induced mutations in the fimH (pilE) gene (ref. 26). At the minimum, it is believed that tests for type 1 fimbriation should include additional functional characterization. While all type 1 fimbriae-mediated adhesion which have been described in this Example is mannose-sensitive, it is not all mannose- or even saccharidespecific as has commonly been thought. Further studies of type 1 fimbriae as a virulence factor must be able to distinguish among the various functional classes.

Allelic variation of the so-called G adhesins of P fimbriated uropathogenic $E.\ coli$ also results in different functional classes, but the requirement for the Gal α l-4Gal sequence within isoreceptors is maintained (refs. 52, 53). These

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differences in G adhesin receptor specificity appear to be rather subtle, at least superficially, when compared to the differences in FimH receptor specificities. Yet there is significantly greater sequence homology among the fimH genes than among the G adhesin genes, some of which share less than 50 percent homology. The G adhesin receptor specificities affect host susceptibility, due in large part to host-specific expression of glycolipid isoreceptor variants. Whether. the FimH family of adhesins bears a similar relationship to 10 host susceptibility or tissue tropism remains to be determined. In this regard it is possible that the G adhesin family could exhibit additional receptor specificities not restricted to the $Gal\alpha l$ -4Gal sequence. The lectin-independent affinity of P fimbriae for immobilized Fn is not dependent on 15 the G adhesin, but on two other minor subunits, E and F, neither of which bear significant homology to FimH (refs. 56, 57).

It is important to point out that the degree of functional heterogeneity of type 1 fimbriae described in the present 20 Example was not appreciated when any of the studies cited above were performed. The results of these studies have made it clear that structural and functional heterogeneity occurs within the class of adhesive organelles commonly referred to as MS or type 1 fimbriae and that the adhesive diversity will lead to a broader spectrum of receptive surfaces for poten-25 tial colonization. The surprising finding that a FimH family of adhesins exists may prove to be an important step toward unravelling the role(s) type 1 fimbriae may play in the ability of enterobacteria to reach their normal habitat or gain entry into deeper tissues, where devastating effects can 30 occur.

EXAMPLE 2

Expression of type 1 fimbriae in heterologous bacterial species

The fim operon of E. coli comprises a cluster of genes covering about 8 kb of DNA. This operon has been isolated and cloned on plasmids in its entirety. The promoter upstream of the fimA gene is located within an invertible DNA sequence, which in E. coli leads to a switch on/switch off situation for fimbrial synthesis. In one orientation of the invertible 10 sequence the promoter is directed towards the fim genes, and the cell is fimbriated; in the other orientation the promoter is directed in the opposite direction, and the cell is nonfimbriated.

Since the regulation of the switch of the invertible promoter sequence is very complex and involves several genes outside the fim operon it is far from certain that the switching on takes place in other bacteria than the enterics. It was therefore considered necessary to insert a replacement promoter for the expression of the fim genes, and as a model for 20 gene expression in a number of different bacterial species the lac promoter was chosen. This promoter has been shown to be active and regulatable in many bacterial species.

Plasmid pPKL83 is a derivative of pPKL4 (ref. 27) carrying the entire fim operon in pBR322, in which the promoter has been destroyed by inserting a BglII linker in the SnaB site located in the promoter sequence. There is a second BglII site in plasmid pPKL83 upstream of the fim operon (Fig. 6). Plasmid pVLT33 (Fig. 7) is a kanamycin resistant derivative of the broad host range plasmid RSF1010, carrying the $lacI^q$ 30 gene and the tac promoter placed upstream of a multiple cloning site in which a unique BamH1 site is placed. The two plasmids were ligated together after digestion of pPKL83 with BglII and pVLT33 with BamH1. In one orientation (pSM1314), this fusion plasmid will express fimbriae in the presence of

IPTG due to the fusion between the fimbrial genes and the lac promoter.

The correct orientation of the fusion plasmid pSM1314 was verified by transforming it into a strain of E. coli which carries a deletion of the fim operon. Production of fimbriae was assayed in two ways: 1) Cell aggregation with fimbrial antibodies and 2) ELISA assay of whole cells. The former analysis is rather simple: to a small volume (10 μ l) of an outgrown or IPTG-induced culture of the cells to be tested is added a small volume (2 μ l) of antibodies raised against 10 fimbriae, on a glass slide. After mixing the samples, fimbriated cells begin to show cell aggregates which are easily observed directly as clumps or under a microscope. No clumping was observed with cells of the strain with a fim deletion, whereas pSM1314 transformants of this strain showed 15 clearly detectable cell aggregates. The ELISA analysis of whole cells confirmed the aggregation assay. In Table 2 below the readings from this type of assay are presented, and they show quantitatively the occurrence of Fim antigens on the cells as a result of IPTG induction of the pSM1314 carrying 20 strain.

Table 2. Results (duplicate) of ELISA assay for type 1 fimbria expressed by pSM1314 in E. coli AAEC191 (OD492)

AAEC191(pSM1314) 0.145/0.164 25 AAEC191(pSM1314)+IPTG 1.026/1.260 Blank 0.113/0.095

Plasmid pSM1314 also carries a mob site which allows it to be transferred to other gram negative bacteria provided a helper plasmid is introduced. This type of transfer is most easily performed in "triparental" matings in which a donor strain (E. coli carrying pSM1314), a helper strain (E. coli carrying a plasmid with conjugation genes) and a recipient strain carrying a selectable marker not present in any of the two



other strains, are mixed on a plate (directly or on a filter). After some growth (often overnight) this mixture is spread on selective plates with antibiotics that only allow the recipient carrying the desired plasmid to grow and form colonies.

In the present context, the *E. coli* strains MC1000(pSM1314) and MC1000(pRK2013) and (as recipient) *Enterobacter cloacae* strain A50 Nal^r (ref. 67), were mated. This recipient strain is resistant to nalidixic acid. After selection for growth on plates with kanamycin plus nalidixic acid the resulting clones were grown in liquid medium and assayed for the presence of fimbriae in the absence/presence of IPTG. The cell aggregation assay was employed.

This assay showed that fimbriae were produced in the Enterobacter cloacae strain and were present on the cell surface;
however, full repression of expression from the tac promoter
was not obtained, most likely due to an increased escape
synthesis. The results showed that E. coli type 1 fimbriae
may be synthesized and processed correctly for pili formation
on the surfaces of heterologous gram-negative bacterial
species.

The plasmid pSM1314 in *E. coli* HB101 was deposited on 26 January 1994 with DSM, the Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH, (German Collection of Microorganisms and Cell Cultures), Mascheroder Weg 1B, D-38124 Braunschweig, Germany, under the accession number DSM 8922.

EXAMPLE 3

The construction of fimH-fusion genes and the expression of mannose-sensitive FimH fusion proteins

30 Heterologous sequences mimicing the pre-S2 region of the hepatitis B viral surface antigen and a neutralizing epitope

of the cholera toxin B chain were inserted in two different positions in the FimH adhesin of type 1 fimbriae. This was carried out by introduction of restriction site handles (BglII-sites) in the fimH gene, followed by in-frame insertion of heterologous DNA segments encoding the foreign epitopes. In the selected positions such insertions did not significantly alter the adhesive function of the FimH protein, since hosts producing hybrid fimbriae that contained the chimeric adhesins exhibited adhesion phenotypes and were 10 normally fimbriated. The heterologous inserts of 52 and 15 amino acids, respectively, residing in the chimeric FimH proteins were recognized by specific sera on the surface of the fimbriae on bacterial hosts. The results illustrate the possibility of using bacterial adhesins as general presenters 15 of foreign antigens and epitopes.

3.1. Materials and methods

3.1.1. Bacterial strain and growth conditions

The Escherichia coli K12 strain HB101 was used in this study as a host for expression of chimeric fimbriae. This strain is phenotypically Fim due to a deletion in the fim gene cluster (ref. 8). Cells were grown on solid medium or in liquid broth supplemented with appropriate antibiotics. When required, gene expression from the lac promoter, residing in front of the fimH gene in plasmid pLPA22 and its derivatives, was induced by the addition of IPTG (isopropyl thiogalactopyranoside) to the growth medium.

3.1.2. Plasmids

Plasmids pPKL4 (comprising the entire, functional fim gene cluster) and pPKL114 (comprising the fimH gene) have been described previously.

pPKL115 which is a plasmid containing the entire type 1 fim gene cluster with a stop linker insertion in the fimH gene

- (i.e. this plasmid expresses all the proteins necessary for the production of fimbriae except the FimH protein) was constructed in two steps:
- (i) plasmid pPKL4 (refs. 27, 28) was digested with KpnI which recognizes a unique restriction site in the fimH gene. The staggered end of the linearized plasmid was made blunt and ligated with the synthetic piece of DNA below (SEQ ID NO:7) containing stop codons in all three reading frames, resulting in plasmid pPKL114:
- 10 5'-GTCGACTTAATTAAGTCGAC-3'
 - 3'-CAGCTGAATTAATTCAGCTG-5';
- (ii) the HindIII-EagI fragment from pPKL114, containing the entire fim gene cluster with the inactivated fimH gene was subsequently inserted into the HindIII and EagI sites of plasmid pACYC184, resulting in plasmid pPKL115.

Plasmid pSM782 (generously provided by S. Molin, Department of Microbiology, Technical University of Denmark, DK-2800 Lyngby) containing the pre-S2 and S encoding regions of the hepatitis B viral genome, was made from plasmid λ -HBV1 (ref. 72) by subcloning a EcoRI-DraI fragment into pBR322.

Plasmid pLPA22 was constructed by inserting a 1018 bp PvuII-MluI fragment containing the fimH gene from pPKL4 into plasmid pUC18. The insert was positioned downstream and in a expression compatible orientation to the lac promoter residing on the vector part of the plasmid (Fig. 8). Expression in E. coli HB101 cells of functional FimH protein was monitored by complementing pLPA22 in trans with pPKL115 and testing for MS adhesion upon induction with IPTG.

Plasmids pLPA29 and pLPA30 were made by inserting 9-mer
30 asymmetric BglII-linkers into the BsaAI and HincII sites,
respectively, in the fimH gene of plasmid pLPA22. At six
different positions in the pLPA22 fimH gene a BglII site was

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introduced without changing the reading frame, resulting in plasmids pLPA98, pLPA36, pLPA58, pLPA30, pLPA29 and pLPA59 (Fig. 10). This was done either by inserting a *BglII* linker into an appropriately treated restriction enzyme site, or by changing 1-3 basepairs using PCR and thereby creating a *BglII* site.

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The plasmid pLPA36 was prepared by opening the pLPA22 fimH gene with the restriction enzyme *Tth1111* and making the ends blunt using Klenow polymerase and ligating using an 8 mer *Bg111* linker (SEQ ID NO:8):

5'-CAGATCTG-3'

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3'-GTCTAGAC-5'

Plasmids pLPA58 and pLPA59 were made by BglII site-creating site-directed mutagenesis of pLPA22 using standard PCR and plasmid pLPA98 was constructed by opening the fimH gene, making the ends blunt with T4 DNA polymerase and ligating with the below 10 mer BglII linker (SEQ ID NO:9):

- 5'-GAAGATCTTC-3'
- 3'-CTTCTAGAAG-5'
- 20 Of the six resulting mutated fimH genes, three expressed protein that was integrated into type 1 fimbriae, and at the same time exhibited mannose-sensitive adhesion. Of these three mutated FimH proteins, the two that conferred to E. coli cells the strongest mannose-sensitive adhesion were expressed from plasmids pLPA29 and pLPA30 (Fig. 9) and these two plasmids were investigated further for their ability to contain large mutations and still be biological active.

Plasmid pLPA29 has a 9 bp long symmetrical BglII linker inserted into the BsaAI site 66 bp upstream of the stop codon for the fimH gene, while plasmid pLPA30 has the same 9 bp BglII linker inserted into the HincII site 163 bp upstream of the stop codon of the fimH gene.

The plasmids pLPA37 and pLPA38 (Fig. 8) were constructed by inserting a 162 bp DNA fragment encoding the pre-S2 region of the Hepatitis B virus surface antigen into the BglII sites in pLPA29 and pLPA30, respectively. This DNA fragment was created by a standard polymerase chain reaction (PCR) using the synthetic primers: (i) 5'-GGAGATCTAATTCCACAACCTT-3' (SEQ ID NO:11) and (ii) 5'-GGAGATCTGTTCAGCGCAGGGT-3' (SEQ ID NO:12), and plasmid pSM782 as a template.

A fragment of plasmid pLPA38 comprising the inserted

heterologous sequence encoding the pre-S2 region of hepatitis
B surface antigen is shown in the below table wherein the
heterologous sequence is underlined and the numbers indicated
correspond to the positions of the amino acid residues in the
mature FimH protein.

Bg/ II
CAG TTC AGA TCT AAT TCC ACA ACC TTC CAC CAA ACT CTG CAA GAT
Gln Phe Arg Ser Asn Ser Thr Thr Phe His Gln Thr Leu Gln Asp
224

CCC AGA GTG AGA GGC CTG TAT TTC CCT GCT GGT GGC TCC AGT TCA Pro Arg Val Arg Gly Leu Tyr Phe Pro Ala Gly Gly Ser Ser Ser

GGA ACA GTA AAC CCT GTT CTG ACT ACT GCC TCT CCC TTA TCG TCA Gly Thr Val Asn Pro Val Leu Thr Thr Ala Ser Pro Leu Ser Ser

ATC TTC TCG AGG ATT GGG GAC CCT GCG CTG AAC AGA TCT TCG ACG
lie Phe Ser Arg lie Gly Asp Pro Ala Leu Asn Arg Ser Ser Thr

226

The plasmids pLPA95 and pLPA93 (Fig. 8) were then made by inserting the below 51 bp synthetic double stranded DNA segment encoding amino acids 50-64 (comprising an epitope) of the B subunit of the cholera toxin into the BglI sites on pLPA30 and pLPA29, respectively (SEQ ID NO:10):

- 5'-GATCTGTTGAAGTTCCGGGTAGTCAGCATATCGATAGTCAGAAAAAAGCTG -3
- 3'- ACAACTTCAAGGCCCATCAGTCGTATAGCTATCAGTCTTTTTTCGACCTAG-5'

A fragment of plasmid pLPA93 comprising the heterologous sequence encoding the above DNA segment of the B subunit of the cholera toxin is shown in the below table wherein the heterologous sequence is underlined and the numbers indicated correspond to the positions of the amino acid residues in the mature FimH protein.

Bg/ II
CAG TTC AGA TCT GTT GAA GTT CCG GGT AGT CAG CAT ATC GAT AGT
GIn Phe Arg Ser Val Glu Val Pro Gly Ser Gln His Ile Asp Ser
224

BamH I/Bg/ II
CAG AAA AAA GCT GGA TCT TCG ACG
GIn Lys Lys Ala Gly Ser Ser Thr
226

3.1.3. DNA techniques

Isolation of plasmid DNA was carried out according to the method of Birnboim and Doly (ref. 73). Restriction endonucleases were used according to the manufacturer's specifications (Biolabs). DNA sequencing was carried out by the dideoxy chain termination technique (ref. 49) using a sequenase version 2.0 kit from USB. Oligonucleotides were made at the core facilities of the Department of Microbiology, Technical University of Denmark.

3.1.4. PCR methodology

Polymerase chain reactions (PCR) were performed on a Perkin 20 Elmer Cetus DNA Thermal Cycler 480. Reactions were set up as 100 μ l volumes containing 200 μ M each of dATP, dCTP, dGTP and dTTP, 0.2-1.0 μ M of each of the two primers, 2 mM MgCl₂, 10 mM Tris-HCl (pH 8.3), 50 mM KCl, 2.5 units of AmpliTaq DNA

polymerase and 0.1-0.2 μg of plasmid template. The reactions were run for 25-30 cycles each consisting of 1 min. at 94°C, 1 min. at 40°C, and 1 min. at 72°C. For amplification of the pre-S2 fragment the above primers 5'GGAGATCTAATTCCACAACCTT 3' (SEQ ID NO:11) and 5'GGAGATCTGTTCAGCGCAGGGT 3' (SEQ ID NO:12) were used.

3.1.5. Hemagglutination

The capacity of bacteria to express a D-mannose binding phenotype was assayed by their ability to agglutinate guinea pig erythrocytes on glass slides. Aliquots of liquid cultures grown to an optical density of 3.0 and 5% erythrocytes were mixed, and the time until agglutination occurred was measured.

3.1.6. Antisera

15 Rabbit anti-type 1 fimbria serum raised against purified type 1 fimbriae has previously been described (ref. 74). A monoclonal antibody directed against FimH (ref. 75) was kindly provided from Dr. Maryvonne Dho-Moulin, Institut National de la Recherche Agronomique, France. Goat serum 20 raised against cholera toxin B subunit (international standard for WHO No. 12-246) produced at the State Serum Institute, Copenhagen, Denmark was kindly provided by same institute. A monoclonal antibody directed against the pre-S2 domain of Hepatitis B surface antigen (ref. 76) was kindly provided by Dr. Makoto Mayumi, Jichi Medical School, Japan. Fluorescein (FITC) conjugated anti rabbit, anti mouse, or anti goat sera were provided from Dakopats, Denmark.

3.1.7. Fluorescence labelling and CCD microscopy

Cells from overnight cultures (IPTG-induced, if required) were harvested, washed in PBS and fixed for 10 minutes at room temperature in a 3.5% (w/v) solution of paraformaldehyde in PBS. Samples of 20 μ l were placed on a poly-L-lysine

coated slide and air dried. After washing in PBS, 16 μ l of a 1:5 (monoclonal) or 1:25 (polyclonal) dilution of the primary antiserum was placed on top of each sample and left in a moist incubation chamber for 1 hour. The slides were washed three times in PBS and 16 μ l of FITC conjugated antiserum were added. After two hours in the dark, the slides were washed three times in PBS and a drop of Citiflour (Citiflour Ltd., London, U.K.) was placed on top of each sample. For visualization, a Carl Zeiss Axioplan microscope equipped for epifluorescence and phase-contrast was employed. Using a charge-coupled device (CCD) camera, pictures were captured as 12-bit files with PMIS software (Photometrics) and subsequently transferred to a Macintosh Quadra 950 computer

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15 <u>3.1.8. Electron microscopy</u>.

for image analysis.

Electron microscopy and immuno-electron microscopy was carried out essentially as described previously (ref. 61). In brief, a 25 μ l aliquot of bacterial suspension was placed on a carbon-coated, glow discharged grid for 30 seconds. Grids were washed in 2 drops of PBS, dehydrated for 5 min in each of the following concentrations of ethanol: 25%, 50%, 75% and 96%, blotted dry and shadowed with tungsten wire at an angle of 30°. For immuno-electron microscopy a monoclonal antibody directed against the pre-S2 region was used diluted 1:5 as the primary antibody and rabbit anti-mouse serum conjugated with 10 nm gold particles (Dako) was used in dilution 1:20 as the secondary antibody.

3.2. Results

As described above, two positions in the C-terminal part of the FimH protein were engineered to contain heterologous sequences mimicing foreign antigenic determinants. In the present study, double plasmid systems were used. In each

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plasmid pair one encoded either a wild-type or an engineered version of the fimH gene, whereas the second plasmid encoded auxiliary functions such as the two-component Fim-specific transport system, regulatory genes and other structural components of the fimbrial organelle except FimH (Table 3).

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3.2.1. Engineering new restriction sites into fimH.

Based on algorithms for prediction of such parameters as hydrophilicity and secondary structure, two potentially optimal positions for insertions of heterologous sequences in 10 the C-terminal domain of the FimH protein were selected. These correspond to positions 225 and 258 in the mature protein predicted to be situated in a surface-exposed part of the FimH protein. In order to facilitate later manipulations, the fimH gene was subcloned into the pUC18 vector resulting 15 in plasmid pLPA22. Subsequently a BglII site was introduced in-frame into positions 225 and 258, respectively. This was carried out by site-directed mutagenesis employing synthetic oligomers resulting in plasmids pLPA30 and pLPA29, respectively (Fig. 9). 20

The introduced BglII sites resulted in a codon change from a Leu to a Phe codon in position 225 and addition of codons for the sequence Arg-Ser-Ser, in the case of plasmid pLPA29, and addition of codons for the sequence Arg-Ser-Gly after position 258 in the case of plasmid pLPA30. Sequence analysis of the entire modified fimH genes in plasmids pLPA29 and pLPA30 confirmed that no other changes had occurred. Host cells which in addition to plasmid pLPA29 or pLPA30 also contained plasmid pPKL115 (fimH), showed wild-type phenotypic characteristics with regard to adhesion and fimbriation as judged by such criteria as hemagglutination (Table 3) and immunofluorescence microscopy.

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3.2.3. Engineering heterologous DNA-sequences encoding the pre-S2 domain of hepatitis B surface antigen and a cholera toxin epitope into fimH.

As heterologous reporter epitopes the pre-S2 region of the

5 hepatitis B surface antigen and a well characterized region
of the B subunit of cholera toxin were selected. The pre-S2
region is known to contain immunologically important (and
protective) antigenic determinants (ref. 76). In addition,
this region is disulphide bond-independent and apparently

10 more immunogenic than the major S protein. The cholera toxin
segment consists of residues 50-64 of the B subunit and has
previously been shown to elicit antibodies that bind to and
neutralize cholera toxin (ref. 77).

A DNA segment of 162 nucleotides encoding 52 of the 55 amino acids of the pre-S2 region was amplified by PCR technology using plasmid pSM782 as template and primers that provided the amplified sequence with flanking BglII sites. Following restriction with BglII and purification the amplified fragment was inserted into the BglII sites of plasmids pLPA29 and pLPA30 resulting in plasmids pLPA37 and pLPA38, respectively (Fig 9). Subsequent sequence analysis confirmed that the inserts were correctly oriented and that the reading frame of the chimeric fimH-pre-S2 genes was correct.

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A synthetic DNA segment encoding the cholera epitope was made by annealing two complementary 51 bp oligonucleotides which were designed to result in a double stranded DNA fragment with a BglII overhang in one end, a BamHI overhang in the other and an internal ClaI site. The epitope-encoding segment was inserted into the BglII site in the fimH gene in plasmids pLPA29 and pLPA30, resulting in regeneration of a BglII site at only one end of the insert. This feature was used to identify plasmids with correct orientation of the insert. The presence of the ClaI site was used for initial screening for clones containing the insert. Sequence analysis of plasmid pLPA93 and pLPA95, both harbouring the epitope-encoding

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segment confirmed the orientation and conservation of the reading frame in the chimeric fimH-cholera genes (Fig 8).

3.2.4. Expression of chimeric FimH adhesin comprising as heterologous sequences the pre-S2 domain of hepatitis B surface antiqen and a cholera toxin epitope.

To evaluate whether the heterologous inserts in fimH were compatible with protein expression the T7 polymerase/promoter system of Tabor and Richardson (ref. 78) was used. Subcloning into the pGEM3 vector system and subsequent assaying revealed that proteins with the expected sizes were produced in all cases from the chimeric fimH genes. More importantly, to assess whether the FimH proteins harbouring foreign inserts were accepted by the type 1 fimbrial transport system and additionally, whether they were present on the bacterial surface in a biologically functional form, the adhesion phenotype of recombinant strains expressing the chimeric FimH proteins was studied.

Bacterial hosts which in addition to plasmid pLPA38 (pre-S2 insert in position 225 in FimH) also contained plasmid pPKL115 (fimH) gave, when induced by IPTG, good agglutination of guinea-pig erythrocytes indicating the presence of a biologically active form of the FimH adhesin on the cells (Table 3). The combination of plasmids pLPA37 (pre-S2 in position 258 in FimH) and pPKL115 resulted in weaker, but detectable, hemagglutination (Table 3). Furthermore, such cells were also shown by electron microscopy to have essentially normal fimbriation (Fig. 10).

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Table 3. Genotype and phenotype of plasmids (A, B or U, respectively indicate pACYC184, pBR322 or pUC18 based vector) used in this study, position of inserts and hemagglutination titer. Host cell: E. coli HB101

5				
	Plasmid	relevant genotype	insert	hemagglu-
			position	tination ^a
	pPKL4 (B)	all <i>fim</i> genes		15
	pPKL115 (A)	fimH		>600
10	pLPA22 (U)	fimH ⁺		>600
	pLPA29 (U)	fimH-BglII	258	>600
	pLPA30 (U)	fimH-BglII	225	>600
	pLPA37 (U)	fimH-pre-S2	258	>600
	pLPA38 (U)	fimH-pre-S2	225	>600
15	pLPA93 (U)	fimH-cholera	225	>600
	pLPA95 (U)	fimH-cholera	258	>600
	pLPA22 (U) +pPKL115 (A)	fimH ⁺ fimH		10
20	pLPA29 +pPKL115	fimH-BglII fimH		7
	pLPA30 +pPKL115	fimH-BglII fimH		8
	pLPA37 +pPKL115	fimH-pre-S2 fimH		210
25	pLPA38 +pPKL115	fimH-pre-S2 fimH		100
	pLPA93 +pPKL115	fimH-cholera fimH		11

fimH-cholera

fimH

30

pLPA95

+pPKL115

a) Hemagglutination of guinea-pig erythrocytes indicated in seconds before reaction occurred. The average values of 4 measurements are given.

In the cases where a sequence mimicing a cholera epitope had been inserted into FimH, viz. pLPA93 (insert in position 225)

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and pLPA95 (insert in position 258), respectively, an agglutination phenotype also resulted when either of these plasmids were complemented by plasmid pPKL115 (fimH) (Table 3). Again, this suggested that in spite of the presence of foreign peptide segments the chimeric FimH proteins were still able to reach the bacterial surface and maintain its adhesive function. In addition to the adherence phenotypes of the various clones the presence of engineered FimH adhesins on the surface of the cells were monitored by CCD microscopy in connection with fluorescent antibody methodology employing a FimH-specific monoclonal serum. In all cases, significant signals, albeit of varying intensity, were detected when compared to a negative control strain that harboured the auxiliary plasmid, pPKL115, alone.

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3.2.5. Immunological detection of the pre-S2 segment of hepatitis B surface antigen and the cholera toxin epitope in chimeric FimH adhesins.

Since there was good evidence that the chimeric FimH proteins were present on the surface of the E. coli hosts the ability of specific antisera, raised against the pre-S2 part of 20 hepatitis B surface antigen or the cholera toxin B chain, respectively to recognize the chimeric FimH-pre-S2 and FimHcholera proteins directly on the surface of the recombinant bacteria were tested. By immunofluorescence microscopy E. 25 coli hosts harbouring either of plasmids pLPA37 or pLPA38 in addition to plasmid pPKL115 were shown to react specifically with antisera directed against the inserted heterologous sequence, whereas hosts expressing wild-type FimH did not. Similar results were obtained with the cholera toxin insert 30 in the same positions (plasmids pLPA93/pPKL115 and pLPA95/pPKL115). Again, the heterologous inserts in the chimeric FimH proteins were recognized by insert-specific serum on the bacterial surface, whereas the relevant control did not react.

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These findings demonstrate that the foreign epitopes are exposed on the surface of extracellularily located chimeric FimH proteins and, significantly, in a conformation which mimics the natural conformation of the epitope(s) as it appears in the native hepatitis B surface antigen or the native cholera toxin.

The results obtained by immunofluorescence microscopy were corroborated by immuno-electron microscopy, employing the pre-S2 specific monoclonal antibody as primary serum and a colloid gold-labelled secondary antiserum. A significant amount of gold particles were seen, mostly in connection with the fimbrial organelles, on bacterial hosts harbouring chimeric fimH-pre-S2 genes (Fig. 10b and 10c), whereas only few goldparticles were present on the control strain expressing wild-type fimbriae (Fig. 10a). Furthermore, in the latter case the gold-particles were not seen to be associated with the fimbriae.

The plasmids pLPA22, pLPA29, pLPA30, pLPA37, pLPA38, pLPA93, pLPA95 and pPKL115 in *E. coli* HB101 were deposited on 26

20 January 1994 with DSM, the Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH, (German Collection of Microorganisms and Cell Cultures), Mascheroder Weg 1B, D-38124 Braunschweig, Germany, under the accession numbers DSM 8915, DSM 8916, DSM 8917, DSM 8918, DSM 8919, DSM 8920, DSM 8921 and DSM 8923, respectively.

EXAMPLE 4

Binding of the MFP class adhesin of E. coli CSH-50 to synthetic peptides

14 synthetic peptides were synthesized on an ABI automated 30 peptide synthesizer according to the method of Merrifield (Merrifield, R.B. 1963. Solid phase peptide synthesis.I. The synthesis of tetrapeptide. J. Am. Chem. Soc. 85:2149). The WO 95/20657 PCT/DK95/00042

binding of the *E. coli* strain CSH-50 to these peptides were tested essentially as described in Example 1. The results of these binding assays indicated that this MFP class strain adhered strongly to one group of peptides whereas the binding the an other group of peptides was absent or weak. In the below listing the one-letter code sequences of the synthetic peptides are shown in a + group, i.e. the group of peptides to which the tested strain adhered strongly, and a - group of peptides to which the binding was weak or absent:

10 + group of peptides

FnSP1: EAQQMVQPQSPVAVSQSKPGCYDNGKHYQI (SEQ ID NO:13)

CB-II-G: EEGKRGARGEBGAAGPVGPBGERGARGNR (SEQ ID NO:14)

sM1(19-32): AIQNIRLRHENKDL (SEQ ID NO:15)

sM6(1-11): RVFPRGTVENPC (SEQ ID NO:16)

15 sM12(1-12): DHSDLVAEKQRLC (SEQ ID NO:17)

sM12(7-18): AEKQRLEDLGQKC (SEQ ID NO:18)

sM5(175-184): TVKDKLAKEQC (SEQ ID NO:19)

sM5(28-54): KTKNEGLKTENEGLKTENEGLKTENEGC (SEQ ID NO:20)

- group of peptides

20 sM5(134-163): QESKENEKALNELLEKTVKDKIAKEQENKE (SEQ ID NO:21)

sM5(117-146): DLTKELNKTRQELANKQQESKENEKALNEL (SEQ ID NO:22)

sM5(14-26): KEALDKYELENHD (SEQ ID NO:23)

sM6(22-31): DVENSMLQAN (SEQ ID NO:24)

sM5(55-84): LKTEKSNLERKTAELTSEKKEHEAENDKLKC (SEQ ID NO:25)

25 sM24(289-303):HQKLEEQNKTSEASRC (SEQ ID NO:26)

EXAMPLE 5

FimH adhesin of further clinical isolates

The following clinical isolates of *E. coli* were tested for adhesion class according to the methods described in Example 1: KB-23, KS-54, U221-3, MJ#9-3, MJ#31-3, MJ#11-2, MJ#2-2.

The results of these experiments are illustrated in Fig. 5. As explained above, the isolate KB-23 showed the M^L type of adhesion, and the isolate U221-3 expressed a M class adhesin showing a mannose-resistant type of adhesion and accordingly, this strain was classified as having a M^R class adhesin. The amino acid sequences of these clinical isolates are shown in Fig. 5 and their nucleotide sequences in Table 5 below.

Table 5 shows the nucleotide sequences of the fimH genes of selected fimH genes disclosed in Example 1 [CI#3 (SEQ ID NO:50), CI#4 (SEQ ID NO:44), CI#7 (SEQ ID NO:51), CI#10 (SEQ ID NO:48) and CI#12 (SEQ ID NO:54)] and as the reference that of the E. coli K12 strain PC31 as it was originally disclosed by Klemm et al. (ref. 27) as the top sequence designated PC31a and the sequence as it was determined recently (PC31b). Additionally, the nucleotide sequences of the following clinical isolates of E. coli are shown: KS54 (SEQ ID NO:52), U221-3 (SEQ ID NO:53), MJ#9-3 (SEQ ID NO:46), MJ#31-3 (SEQ ID NO:47), MJ#11-2 (SEQ ID NO:43), MJ#2-2 (SEQ ID NO:45) and F-18 (SEQ ID NO:42).

Table 5. Nucleotide sequences of the above fimH genes disclosed in Example 1, E. coli K12 strain PC31 (PC31a and PC31b) and the nucleotide sequences of KS54, U221-3, MJ#9-3, MJ#31-3, MJ#11-2, MJ#2-2 and F-18.

	109	14																
PC31a		-	CCR	C 75 75	1.00	100	~==											
F-18	770	AAA	CGA	GII	ATT	ACC	CTG	TTT	GCT	GTA	CTG	CTG	ATG	GGC	TGG	TCG	GTA	AAT
MJ11-2																		
CI 4																		
MJ22																		
MJ9-3																		
MJ31-3																		
CI 10																		
PC31b																		
CI 3																		
CI 7						-A-												
KS54						-A-												
U221-3																		
CI 12																		
C1 12																		
	114	0																
PC31a		•																
F-18		TGG	TÇA	TTC	GCC	TGT	AAA	ACC	GCC	AAT	GGT	ACC	GCT	ATC	CCT	ATT	GGC	GGT
MJ11-2													A					
CI 4													A					
MJ22														-,				
MJ9-3													X					
MJ31-3														T				
CI 10														T				
PC31b																		
CI 3																		
CI 7											c							
KS54											c							
U221-3																		
CI 12																		
	1202	,																
PC31a		AGC	GCC	AAT	GTT	ቸልሞ	GT N	220										
F-18										GCG	_ ~	GIC	GTG	AAT	GTG	GGG	CAA	AAC
MJ11-2												^						
CI 4											T							
MJ22											T	-C-						
MJ9-3			T								m							
MJ31-3			T								T							
CI 10											T	-C-						
												-0-						
CI 3																		
CI 7									- 			-C-						
KS54			Т									-0-						C
U221-3									-		T	-C-						
CI 12									- 			-C-						

Table 5, continued

PC31a	CTC	GTC	GTG	GAT		TCG	ACC	- CB B										
F-18							700	CAA	ATC	TTT	TGC	CAT	AAC	GAT	TAT	CCG	GAA	ACC
MJ11-2			A												·c	A		
CI 4															·c	A		
MJ22			4												·c			
MJ9-3			A												c	A		
MJ31-3			A															
CI 10																		
PC31b																		
CI 3																		
CI 7			3												C			
KS54			A												C			
U221-3																		
CI 12																		
															C			
	131	2							,									
PC31a	ATT	ACA	GAC	TAT	GTC	ACA	CTG	CAA	CCN		***	GCT						
F-18																		
MJ11-2										~~								
CI 4										T								
MJ22										T			_		-			
MJ9-3																		
MJ31-3																		
CI 10																		
PC31P																		
CI 3			T										_					
CI 7										T								
KS54																		
U221-3																		
CI 12										T								
	1366	-																
PC31a	AAT	TTI	TCC	GGG	ACC	GTA	AAA	TAT	AGT	GGC	AGT	AGC	TAT	CCA	TTT	CCT	ACC	ACC
F-18 MJ11-2	-								-4-					-	_			
MJ11-2 CI 4	•								- A -					-	_			
MJ22	- 0-			-4-					- A -					_	_			
MJ9-3									•									
MJ31-3																		
CI 10																		
CI 3																		
CI 7					- .											G		
KS54					,	·		~~~ .										
U221-3						- .										G	T	
CI 12	-G-				:	- .	- 											
	•		- '	'					-A-					T	c ·		T	

Table 5, continued

	4 44																	
PC31a	AG	C GA	A AC	G CC	G CGC	GTI	GTI	TAT		* ***								
F-18					G				AAI	100	3 AGA	L ACG	GAT	AAG	CCG	TGG	CCG	GTG
MJ11-2					G													
CI 4					G													
MJ22																		
MJ9-3					G													
MJ31-3																		
CI 10																		
PC31b																		
CI 3																		
CI 7				- T	·G													
KS54																		
U221-3																		
CI 12																		
	147	-																
PC31a	GCG	CTI	TAT	TTG	ACG	CCT	GTG	ACC	ACT	~~~								
F-18						G			AGI		GGC	GGG	GIG	GCG	ATT	AAA	GCT	GGC
MJ11-2																		
CI 4																		
MJ22																		
MJ9-3																		
MJ31-3											T							
CI 10																		
PC31b															-			
CI 3																		
CI 7																		
KS54											~							
U221-3		-G-									T							
CI 12																		
											G	~-A						
	152																	
PC31a			2 000															
F-18		117	MII	GCC	GTG	CTT	ATT	TTG	CGA	CAG	ACC	AAC	AAC	TAT .	AAC .	AGC (GAT (GAT
MJ11-2																		
CI 4																		
MJ22																		
MJ9-3																		
MJ31-3																		
CI 10																		
PC31b																		
CI 3																		
CI 7																		
KS54																		
U221-3																		
CI 12								~			- .							

Table 5, continued

PC31a	T	rc d	CAG	TT	r ca	rc	~~			_												
F-18							GG	AA	AT	T TA	CG	CC A	AT	AAT	GA:	r GTC	GTO	G GT	C C	~~ *	~~	
MJ11-	.2	:														GTC			- C	-1 N	CT	GG
MJ11- CI 4																						
MJ22																				-c -		
																				-c -		
MJ9-3																				-c -		
MJ31- CI 10	3							_														
CI 10																						
PC31b							'															
CI 3																						
CI 7																						
KS54																				·	-	
U221-:	a																				-	
CI 12	,																					
C1 12																						
																				c		
	163																					
PC31a	GGC	T	3C (GAT	GTT	*	~ ~	~~														
F-18							- 0	rC1	CGT	GAT	GI	C AC	C	STT	ACT	CTG	CCG	GAC	TAC		T (~
MJ11-2			·т -		-6-											CTG						-31
CI 4			· T		-0-																	
MJ22			· T													T				· -c.		
MJ9-3					C											T				c.		
MJ31-3									-A-											-C-		
									-A-											-C-		
CI 10									G											-c-		
PC31b																				-C-		
CI 3																				-c-	-	
CI 7																				-C-	-	
KS54									_					~						-0-		
U221-3							_		••				-							-0-		
CI 12			T -					_					-							-CA		
								-	.~~						'	 r				-C-		
	1690	0																		_		_
PC31a	TCA	GTO		~ n ,				_								_						
F-18				A	411	CCT	CI	T	rcc (GTT	TAT	TGT	GC	G A	AA I	AGC C	'AA 1	24	~~~	~~~		_
MJ11-2				-												AGC C			C16	GGG	TA	.T
CI 4				-G -																		-
MJ22				-G -																		-
MJ9-3				-G -											- 							-
MJ31-3																						-
=																 						-
CI 10																						_
PC31P																						_
CI 3	~			G -																		_
CI 7																·						_
KS54																						-
U221-3																						•
CI 12	:			·				-									-					•
_				·																		•

Table 5, continued

PC31a																		
F-18	TAC	CTC	TCC	GGC	: ACA	CAC	: GC#	GA1	GCG	GGC	AAC	TCG	ATT	TIC	ACC	AAT	ACC	GCG
MJ11-2																		
CI 4		. – –				1												
MJ22						G					~							
MJ9-3		·																
MJ31-3																		
CI 10 PC31b																		
CI 3																		
CI 7												_						
KS54																		_
U221-3																		
CI 12																		
	179	8																
PC31a	TCG	TTT	TCA	CCT	GCA	CAG	GGC	GTC	GGC	GTA	CAG	TTG	ACG	CGC	AAC	GGT	ACG	ልተተ
F-18					6													
MJ11-2				C	G													
CI 4																		
MJ22				C	G													
MJ9-3				A	G													
MJ31-3				A	G													
CI 10											_							
bc37P											_							
CI 3																		
CI 7				A	G													
KS54				A	G													
U221-3																		
CI 12																		
	185																	
PC31a	ATT	CCA	GCG	AAT	AAC	ACG	GTA	TCG	TTA	GGA	GCA	GTA	GGG	ACT	TCG	GCG	GTG	ACT
F-18 MJ11-2																		
CI 4																		
MJ22																		
MJ9-3																		
MJ31-3											_							
CI 10																		
PC31P																		
CI 3																		
CI 7																		
KS54					~													
U221-3																		
CI 12															:		A	

Table 5, continued

	190)6															
PC31a	CTG	GGA	TTA	ACG	GCA	AAT	TAT	GCA	CGT	ACC	GGA	GGG	CNC	CTC			
F-18							C										
MJ11-2																	
CI 4							C										
MJ22							0										
MJ9-3							0								c		
MJ31-3							0										
CI 10	T																
PC31b																	
CI 3																	
CI 7																	
KS54																	
U221-3																	
CI 12																	
CI 12							C										
CI 12	196						- - C										
PC3la	196	0															
	196 GTG	O CAA	TCG	ATT	ATT	GGC	GTG	ACT	TTT	GTT	727	CAA					
PC31a	196 GTG	CAA	TCG	ATT	ATT	GGC	GTG	ACT	TTT	GTT	TAT	CAA					
PC31a F-18	196 GTG	CAA	TCG	ATT	ATT	GGC	GTG	ACT	TTT	GTT	TAT	CAA					
PC31a F-18 MJ11-2	196 GTG	CAA	TCG	ATT	ATT	GGC	GTG	ACT	TTT	GTT	TAT	CAA					
PC31a F-18 MJ11-2 CI 4	196 GTG	CAA	TCG	ATT	ATT	GGC	GTG	ACT	TTT	GTT	TAT	CAA					
PC31a F-18 MJ11-2 CI 4 MJ22	196 GTG	CAA	TCG	ATT	ATT	GGC	GTG	ACT	TTT	GTT	TAT	CAA				•••	
PC31a F-18 MJ11-2 CI 4 MJ22 MJ9-3	196 GTG	CAA	TCG	ATT	ATT	GGC	GTG	ACT	TTT	GTT	TAT	CAA					
PC31a F-18 MJ11-2 CI 4 MJ22 MJ9-3 MJ31-3	196 GTG	CAA	TCG	ATT	ATT	GGC	GTG	ACT	TTT	GTT	TAT	CAA					
PC31a F-18 MJ11-2 CI 4 MJ22 MJ9-3 MJ31-3 CI 10	196 GTG	CAA	TCG	ATT	ATT	GGC	GTG	ACT	TTT	GTT	TAT	CAA					
PC31a F-18 MJ11-2 CI 4 MJ22 MJ9-3 MJ31-3 CI 10 PC31b	196 GTG	CAA	TCG	ATT	ATT	GGC	GTG	ACT	TTT	GTT	TAT	CAA					
PC31a F-18 MJ11-2 CI 4 MJ22 MJ9-3 MJ31-3 CI 10 PC31b CI 3	196 GTG	CAA	TCG	ATT	ATT	GGC	GTG	ACT	TTT	GTT	TAT	CAA					
PC31a F-18 MJ11-2 CI 4 MJ22 MJ9-3 MJ31-3 CI 10 PC31b CI 3 CI 7 KS54	196 GTG	CAA	TCG	ATT	ATT	GGC	GTG	ACT	TTT	GTT	TAT	CAA					

EXAMPLE 6

Enrichment selection of strains having mutated FimH adhesins conferring altered adhesion ability

One mechanism whereby new binding activities of bacterial
adhesins may arise is by random, naturally occurring mutagenesis. In nature, a variety of factors would enrich for strains that possessed adhesive capacities conferring a selective advantage. In the present example an in vitro procedure was used to select for potential mutants with altered adhesive capacity. As a target substratum bovine κ-casein was selected.

 κ -casein is the glycosylated isoform of bovine casein consists of a single polypeptide chain containing 169 amino acid residues the sequence of which has been determined (ref. 68). Bovine κ -casein does not contain N-glycosidic linkages, but 15 up to six O-linked oligosaccharides are present in the Cterminal region of the molecule (refs. 68, 69). The saccharide moieties are heterologous and also vary as a function of time after parturition. Of significance for the present study is the fact that D-mannose is not present in the bovine κ -20 casein. Only di- to hexasaccharides containing galactose, Nacetyl-galactosamine, N-acetyl-glucosamine, fucose and sialic acid have been described (ref. 68). Glycoproteins having such saccharide compositions would not be expected to serve as a receptor for the classic type of the FimH adhesin such as is 25 found in E. coli strain PC31.

Adhesion tests were performed to verify the inability of recombinant strains carrying the fimH gene from E. colistrain PC31 to adhere to immobilized κ -casein. The E. colistrain used, KB1001 is HB101 containing plasmids pPKL115 and pLPA22 (ref. 70). The adhesion assay was performed using microtiter plates coated with 30 μ g/ml κ -casein in 0.1 M sodium bicarbonate (pH 9.6) for 30 minutes, followed by blocking any remaining binding sites with a subsequent in-

cubation with 0.1% bovine serum albumin (BSA) in PBS. A quantitative adhesin assay was performed as described in more detail elsewhere (ref. 71). Briefly, bacteria were diluted to equivalent concentrations (5 x 10⁷ cells/100µl) in PBS

5 containing 0.1% BSA, added to coated microtiter wells for 30 minutes at 37°C. After washing the wells thoroughly to remove unbound bacteria, BHI broth was added and the bacteria were allowed to grow at 37°C on a rotating platform (150 rpm) until the optical density could be measured (2-2.5 hours).

10 Comparisons can be made of optical densities obtained in the test wells to those obtained in standard curves developed from the plating of known numbers of bacteria under similar conditions, allowing extrapolation to absolute numbers of bound bacteria (ref. 70).

15 The KB1001 strain comprising the fimH gene from PC31 bound to immobilized mannan in significant numbers, but there was substantially no measurable adhesion to immobilized κ -casein. To select for possible mutant cells having acquired the ability to bind to κ -casein, cells of KB1001 were allowed to 20 interact with κ -casein immobilized on microtiter wells. After thorough washing to remove non-adhering bacterial cells, cells adhering to the wells were collected and grown overnight in BHI broth. These "enriched" bacterial cultures were again allowed to interact with immobilized κ -casein, the 25 plates were washed and adhering cells collected in nutrient broth. This enrichment cycle was repeated up to ten times. Bacterial cells obtained from the last of these cycles ("enriched" strains) adhered to k-casein in significantly increased numbers in comparison to the parent ("non-enriched") strain (Table 6.1). Individual colonies of "enriched" KB1001 30 were isolated and four tested for ability to adhere to κ casein. Three enriched cultures (clones) bound to κ -casein significantly better than did the non-enriched parent strain.

Table 6.1. Adhesion to casein of non-enriched and enriched E. coli strain KB1001.

5 Strain

bacteria binding to

κ-casein^{a)}

Non-enriched KB1001

 0.043 ± 0.018

(pPKL115 + pLPA22)

Enriched KB1001

 0.249 ± 0.004

10 (pPKL115 + pLPA22)

a) numbers represent optical density of bacterial growth \pm S.D. with background O.D. substracted. N = 3.

To determine whether the new adhesive activity was due to plasmid-related changes and not simply to host cell-related changes, plasmid preparations of pLPA22 were made from enriched and from non-enriched strains and used to transform E. coli HB101 containing the auxiliary plasmid pPKL115. Randomly selected transformants resistant to ampicillin and chloramphenical were tested for adhesion to κ-casein, and several of the transformants harbouring plasmids from enriched cultures adhered in significantly increased numbers relative to plasmid-containing cells of the non-enriched parent strain (Table 6.2).

Table 6.2. Adhesion to casein of HB101 (pPKL115) transformed with plasmids from enriched or non-enriched strain KB1001.

Plasmid derived from:

bacteria binding to

 κ -casein^{a)}

Non-enriched KB1001

 $5 + 0.1 \times 10^3$

30 Enriched KB1001

 $50 \pm 1.5 \times 10^{3}$

a) numbers represent mean number of bacteria per well \pm S.D. N = 3.

The above results demonstrate that random or spontaneous mutations in genes coding for a bacterial adhesin that confer binding to a new substratum (i.e. a receptor moiety to which the parent strain does not bind), can be selected for by appropriate in vitro procedures.

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INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made below relate to the microorganism ref on page 53, line 26	
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet
Name of depositary institution	
DSM-Deutsche Sammlung von Mikroorg	ganismen und Zellkulturen GmbH
Address of depositary institution (including postal code and country)	
Mascheroder Weg 1B D-38124 Braunschweig Germany	
Date of deposit	Accession Number
26 January 1994	DSM 8922
C. ADDITIONAL INDICATIONS (leave blank if not applicable	(e) This information is continued on an additional sheet
As regards the respective Patent nated states, the applicants required microorganisms only be made a by the requester until the date of the date on which the application is deemed to be withdrawn.	est that a sample of the deposi- vailable to an expert nominated n which the patent is granted or
D. DESIGNATED STATES FOR WHICH INDICATIO	NS ARE MADE (if the indications are not for all designated States)
	·
E. SEPARATE FURNISHING OF INDICATIONS (laws	blank if not applicable)
The indications listed below will be submitted to the International I Number of Deposit?	Bureau later (specify the general nature of the indications e.g., "Accession
For receiving Office use only	For International Bureau use only
This sheet was received with the international application	This sheet was received by the International Bureau on:
Authorized officer Guscur Gelducill	Authorized officer



INDICATIONS RELATING TO DEPOSITED MICROORGANISMS (PCT Rule 12bis)

Additional sheet

In addition to the microorganism indicated on page 53 of the description, the following microorganisms have been deposited with

DSM-Deutsche Sammlung von Mikroorganismen und Cellkulturen GmbH Mascheroder Weg 1b, D-38124 Braunschweig, Germany

10 on the dates and under the accession numbers as stated below:

	Accession number	Date of deposit		Description Page No.	Description Line No.
	DSM 8915	26 January	1994	66	24
15	DSM 8916	26 January	1994	66	24
	DSM 8917	26 January	1994	66	24
	DSM 8918	26 January	1994	66	24
	DSM 8919	26 January	1994	66	24
	DSM 8920	26 January	1994	66	24
20	DSM 8921	26 January		66	25
	DSM 8923	26 January	1994	66	25

For all of the above-identified deposited microorganisms, the following additional indications apply:

As regards the respective Patent Offices of the respective designated states, the applicants request that a sample of the deposited microorganisms stated above only be made available to an expert nominated by the requester until the date on which the patent is granted or the date on which the application has been refused or withdrawn or is deemed to be withdrawn.

SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT:
 - (A) NAME: GX BioSystems A/S
 - (B) STREET: Mothsvej 70
 - (C) CITY: Holte
 - (D) COUNTRY: Denmark
 - (E) POSTAL CODE (ZIP): 2840
- (ii) TITLE OF INVENTION: Receptor specific bacterial adhesins and their use
- (iii) NUMBER OF SEQUENCES: 55
 - (iv) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 300 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:
 - Met Lys Arg Val Ile Thr Leu Phe Ala Val Leu Leu Met Gly Trp Ser

 1 10 15
 - Val Asn Ala Trp Ser Phe Ala Cys Lys Thr Ala Asn Gly Thr Ala Ile 20 25 30
 - Pro Ile Gly Gly Ser Ala Asn Val Tyr Val Asn Leu Ala Pro Val 35 40 45
 - Val Asn Val Gly Gln Asn Leu Val Val Asp Leu Ser Thr Gln Ile Phe 50 55 60
 - Cys His Asn Asp Tyr Pro Glu Thr Ile Thr Asp Tyr Val Thr Leu Gln 65 70 75 80
 - Arg Gly Ser Ala Tyr Gly Gly Val Leu Ser Asn Phe Ser Gly Thr Val 85 90 95
 - Lys Tyr Ser Gly Ser Ser Tyr Pro Phe Pro Thr Thr Ser Glu Thr Pro 100 105 110

Arg Val Val Tyr Asn Ser Arg Thr Asp Lys Pro Trp Pro Val Ala Leu 115 120 125

Tyr Leu Thr Pro Val Ser Ser Ala Gly Gly Val Ala Ile Lys Ala Gly 130 135 140

Ser Leu Ile Ala Val Leu Ile Leu Arg Gln Thr Asn Asn Tyr Asn Ser 145 150 155 160

Asp Asp Phe Gln Phe Val Trp Asn Ile Tyr Ala Asn Asn Asp Val Val
165 170 175

Val Pro Thr Gly Gly Cys Asp Val Ser Ala Arg Asp Val Thr Val Thr 180 185 190

Leu Pro Asp Tyr Pro Gly Ser Val Pro Ile Pro Leu Thr Val Tyr Cys 195 200 205

Ala Lys Ser Gln Asn Leu Gly Tyr Tyr Leu Ser Gly Thr His Ala Asp 210 215 220

Ala Gly Asn Ser Ile Phe Thr Asn Thr Ala Ser Phe Ser Pro Ala Gln 225 230 235 240

Gly Val Gly Val Gln Leu Thr Arg Asn Gly Thr Ile Ile Pro Ala Asn 245 250 255

Asn Thr Val Ser Leu Gly Ala Val Gly Thr Ser Ala Val Ser Leu Gly 260 265 270

Leu Thr Ala Asn Tyr Ala Arg Thr Gly Gly Gln Val Thr Ala Gly Asn 275 280 285

Val Gln Ser Ile Ile Gly Val Thr Phe Val Tyr Gln 290 295 300

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 30 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Glu Ala Gln Gln Met Val Gln Pro Gln Ser Pro Val Ala Val Ser Gln
1 10 15

Ser Lys Pro Gly Cys Tyr Asp Asn Gly Lys His Tyr Gln Ile 20 25 30

(2) INFORMATION FOR SEQ ID NO:3:

	(A) LENGTH: 29 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
GGG	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:	29
	INFORMATION FOR SEQ ID NO:4:	29
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 29 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:	
GGG	GGTGCAC TCAGGGAACC ATTCAGGCA	29
(2)	INFORMATION FOR SEQ ID NO:5:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 30 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:	
GGGT	IGCGCAT TATTGATAAA CAAAAGTCAC	30
(2)	INFORMATION FOR SEQ ID NO:6:	
	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 30 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	(ii) MOLECULE TYPE: DNA (genomic)	

((xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:	
GGGGC	CATGCT TATTGATAAA CAAAAGTCAC	30
(2) I	INFORMATION FOR SEQ ID NO:7:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	
GTCGA	CTTAA TTAATTAAGT CGAC	24
(2) I	NFORMATION FOR SEQ ID NO:8:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 8 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	ii) MOLECULE TYPE: DNA (genomic)	
(.	xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:	
CAGAT	CTG	8
(2) I	NFORMATION FOR SEQ ID NO:9:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 10 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(:	xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:	
GAAGA'	TCTTC	10
(2) I	NFORMATION FOR SEQ ID NO:10:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 51 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	

		(D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:10:	
GAT		TT TTTTCTGACT ATCGATATGC TGACTACCCG GAACTTCAAC A	
			51
(2)	INFC	ORMATION FOR SEQ ID NO:11:	
	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 22 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:11:	
GGAG	ATCT	AA TTCCACAACC TT	22
(2)	INFO	RMATION FOR SEQ ID NO:12:	
	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 22 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:12:	
GGAG	ATCT	GT TCAGCGCAGG GT	22
(2)	INFO	RMATION FOR SEQ ID NO:13:	
	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 30 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: unknown (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: peptide	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:13:	
	Glu 1	Ala Gln Gln Met Val Gln Pro Gln Ser Pro Val Ala Val Ser Gln 5 10 15	

Ser Lys Pro Gly Cys Tyr Asp Asn Gly Lys His Tyr Gln Ile 20 25 30

- (2) INFORMATION FOR SEQ ID NO:14:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 29 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

Glu Glu Gly Lys Arg Gly Ala Arg Gly Glu Asx Gly Ala Ala Gly Pro 1 5 10 15

Val Gly Pro Asx Gly Glu Arg Gly Ala Arg Gly Asn Arg 20 25

- (2) INFORMATION FOR SEQ ID NO:15:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

Ala Ile Gln Asn Ile Arg Leu Arg His Glu Asn Lys Asp Leu 1 5 10

- (2) INFORMATION FOR SEQ ID NO:16:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 12 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

Arg Val Phe Pro Arg Gly Thr Val Glu Asn Pro Cys
1 5 10

(2) INFORMATION FOR SEQ ID NO:17:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 13 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:

Asp His Ser Asp Leu Val Ala Glu Lys Gln Arg Leu Cys
1 10

- (2) INFORMATION FOR SEQ ID NO:18:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 13 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

Ala Glu Lys Gln Arg Leu Glu Asp Leu Gly Gln Lys Cys
1 10

- (2) INFORMATION FOR SEQ ID NO:19:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 11 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

Thr Val Lys Asp Lys Leu Ala Lys Glu Gln Cys
1 5 10

- (2) INFORMATION FOR SEQ ID NO:20:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 28 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

Lys Thr Lys Asn Glu Gly Leu Lys Thr Glu Asn Glu Gly Leu Lys Thr

1 10 15

Glu Asn Glu Gly Leu Lys Thr Glu Asn Glu Gly Cys 20 25

- (2) INFORMATION FOR SEQ ID NO:21:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 30 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:

Gln Glu Ser Lys Glu Asn Glu Lys Ala Leu Asn Glu Leu Leu Glu Lys

1 10 15

Thr Val Lys Asp Lys Ile Ala Lys Glu Glu Asn Lys Glu 20 25 30

- (2) INFORMATION FOR SEQ ID NO:22:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 30 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:

Asp Leu Thr Lys Glu Leu Asn Lys Thr Arg Gln Glu Leu Ala Asn Lys

1 10 15

Gln Gln Glu Ser Lys Glu Asn Glu Lys Ala Leu Asn Glu Leu 20 25 30

- (2) INFORMATION FOR SEQ ID NO:23:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 13 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown

- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:

Lys Glu Ala Leu Asp Lys Tyr Glu Leu Glu Asn His Asp 1 5 10

- (2) INFORMATION FOR SEQ ID NO:24:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 10 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:

Asp Val Glu Asn Ser Met Leu Gln Ala Asn 1 5 10

- (2) INFORMATION FOR SEQ ID NO:25:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 31 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:

Leu Lys Thr Glu Lys Ser Asn Leu Glu Arg Lys Thr Ala Glu Leu Thr 1 5 10 15

Ser Glu Lys Lys Glu His Glu Ala Glu Asn Asp Lys Leu Lys Cys 20 25 30

- (2) INFORMATION FOR SEQ ID NO:26:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 16 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:

His Gln Lys Leu Glu Glu Gln Asn Lys Thr Ser Glu Ala Ser Arg Cys
1 10 15

- (2) INFORMATION FOR SEO ID NO:27:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 300 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:

Met Lys Arg Val Ile Thr Leu Phe Ala Val Leu Leu Met Gly Trp Ser 1 5 10 15

Val Asn Ala Trp Ser Phe Ala Cys Lys Thr Ala Asn Gly Thr Ala Ile 20 25 30

Pro Ile Gly Gly Ser Ala Asn Val Tyr Val Asn Leu Ala Pro Ala 35 40 45

Val Asn Val Gly Gln Asn Leu Val Val Asp Leu Ser Thr Gln Ile Phe 50 55 60

Cys His Asn Asp Tyr Pro Glu Thr Ile Thr Asp Tyr Val Thr Leu Gln 65 70 75 80

Arg Gly Ser Ala Tyr Gly Gly Val Leu Ser Asn Phe Ser Gly Thr Val 85 90 95

Lys Tyr Ser Gly Ser Ser Tyr Pro Phe Pro Thr Thr Ser Glu Thr Pro 100 105 110

Arg Val Val Tyr Asn Ser Arg Thr Asp Lys Pro Trp Pro Val Ala Leu 115 120 125

Tyr Leu Thr Pro Val Ser Ser Ala Gly Gly Val Ala Ile Lys Ala Gly
130 135 140

Ser Leu Ile Ala Val Leu Ile Leu Arg Gln Thr Asn Asn Tyr Asn Ser 145 150 155 160

Asp Asp Phe Gln Phe Val Trp Asn Ile Tyr Ala Asn Asn Asp Val Val 165 170 175

Val Pro Thr Gly Gly Cys Asp Val Ser Ala Arg Asp Val Thr 180 185 190

Leu Pro Asp Tyr Pro Gly Ser Val Pro Ile Pro Leu Thr Val Tyr Cys

200

205

Ala Lys Ser Gln Asn Leu Gly Tyr Tyr Leu Ser Gly Thr His Ala Asp 210 215 220

Ala Gly Asn Ser Ile Phe Thr Asn Thr Ala Ser Phe Ser Pro Ala Gln 225 230 235 240

Gly Val Gly Val Gln Leu Thr Arg Asn Gly Thr Ile Ile Pro Ala Asn 245 250 255

Asn Thr Val Ser Leu Gly Ala Val Gly Thr Ser Ala Val Ser Leu Gly 260 265 270

Leu Thr Ala Asn Tyr Ala Arg Thr Gly Gly Gln Val Thr Ala Gly Asn 275 280 285

Val Gln Ser Ile Ile Gly Val Thr Phe Val Tyr Gln 290 295 300

(2) INFORMATION FOR SEQ ID NO:28:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 300 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:

Met Lys Arg Val Ile Thr Leu Phe Ala Val Leu Leu Met Gly Trp Ser 1 5 10 15

Val Asn Ala Trp Ser Phe Ala Cys Lys Thr Ala Asn Gly Thr Ala Ile 20 25 30

Pro Ile Gly Gly Ser Ala Asn Val Tyr Val Asn Leu Ala Pro Ala 35 40 45

Val Asn Val Gly Gln Asn Leu Val Val Asp Leu Ser Thr Gln Ile Phe 50 55 60

Cys His Asn Asp Tyr Pro Glu Thr Ile Thr Asp Tyr Val Thr Leu Gln 65 70 75 80

Arg Gly Ser Ala Tyr Gly Gly Val Leu Ser Ser Phe Ser Gly Thr Val 85 90 95

Lys Tyr Asn Gly Ser Ser Tyr Pro Phe Pro Thr Thr Ser Glu Thr Pro 100 105 110

Arg Val Val Tyr Asn Ser Arg Thr Asp Lys Pro Trp Pro Val Ala Leu 115 120 125 Tyr Leu Thr Pro Val Ser Ser Ala Gly Gly Val Ala Ile Lys Ala Gly 130 135 140

Ser Leu Ile Ala Val Leu Ile Leu Arg Gln Thr Asn Asn Tyr Asn Ser 145 150 155 160

Asp Asp Phe Gln Phe Val Trp Asn Ile Tyr Ala Asn Asn Asp Val Val 165 170 175

Val Pro Thr Gly Gly Cys Asp Val Ser Ala Cys Asp Val Thr Val Thr 180 185 190

Leu Pro Asp Tyr Pro Gly Ser Val Pro Ile Pro Leu Thr Val Tyr Cys
195 200 205

Ala Lys Ser Gln Asn Leu Gly Tyr Tyr Leu Ser Gly Thr His Ala Asp 210 215 220

Ala Gly Asn Ser Ile Phe Thr Asn Thr Ala Ser Phe Ser Pro Ala Gln 225 230 235 240

Gly Val Gly Val Gln Leu Thr Arg Asn Gly Thr Ile Ile Pro Ala Asn 245 250 255

Asn Thr Val Ser Leu Gly Ala Val Gly Thr Ser Ala Val Ser Leu Gly 260 265 270

Leu Thr Ala Asn Tyr Ala Arg Thr Gly Gly Gln Val Thr Ala Gly Asn 275 280 285

Val Gln Ser Ile Ile Gly Val Thr Phe Val Tyr Gln 290 295 300

(2) INFORMATION FOR SEQ ID NO:29:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 300 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:

Met Lys Arg Val Ile Thr Leu Phe Ala Val Leu Leu Met Gly Trp Ser 1 5 10 15

Val Asn Ala Trp Ser Phe Ala Cys Lys Thr Ala Asn Gly Thr Ala Ile 20 25 30

Pro Ile Gly Gly Ser Ala Asn Val Tyr Val Asn Leu Ala Pro Ala 35 40 45

Val Asn Val Gly Gln Asn Leu Val Val Asp Leu Ser Thr Gln Ile Phe 50 55 60

Cys His Asn Asp Tyr Pro Glu Thr Ile Thr Asp Tyr Val Thr Leu Gln 65 70 75 80

Arg Gly Ser Ala Tyr Gly Gly Val Leu Ser Ser Phe Ser Glu Thr Val 85 90 95

Lys Tyr Asn Gly Ser Ser Tyr Pro Phe Pro Thr Thr Ser Glu Thr Pro 100 105 110

Arg Val Val Tyr Asn Ser Arg Thr Asp Lys Pro Trp Pro Val Ala Leu 115 120 125

Tyr Leu Thr Pro Val Ser Ser Ala Gly Gly Val Ala Ile Lys Ala Gly
130 135 140

Ser Leu Ile Ala Val Leu Ile Leu Arg Gln Thr Asn Asn Tyr Asn Ser 145 150 155 160

Asp Asp Phe Gln Phe Val Trp Asn Ile Tyr Ala Asn Asn Asp Val Val 165 170 175

Val Pro Thr Gly Gly Cys Asp Val Ser Ala Arg Asp Val Thr Val Thr 180 185 190

Leu Pro Asp Tyr Pro Gly Ser Val Pro Ile Pro Leu Thr Val Tyr Cys
195 200 205

Ala Lys Ser Gln Asn Leu Gly Tyr Tyr Leu Ser Gly Thr Asp Ala Asp 210 215 220

Ala Gly Asn Ser Ile Phe Thr Asn Thr Ala Ser Phe Ser Pro Ala Gln 225 230 235 240

Gly Val Gly Val Gln Leu Thr Arg Asn Gly Thr Ile Ile Pro Ala Asn 245 250 255

Asn Thr Val Ser Leu Gly Ala Val Gly Thr Ser Ala Val Ser Leu Gly 260 265 270

Leu Thr Ala Asn Tyr Ala Arg Thr Gly Gly Gln Val Thr Ala Gly Asn 275 280 285

Val Gln Ser Ile Ile Gly Val Thr Phe Val Tyr Gln 290 295 300

(2) INFORMATION FOR SEQ ID NO:30:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 300 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:

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Met Lys Arg Val Ile Asn Leu Phe Ala Val Leu Leu Met Gly Trp Ser 5 10 Val Asn Ala Trp Ser Phe Ala Cys Lys Thr Ala Asn Gly Thr Ala Ile Pro Ile Gly Gly Ser Ala Asn Val Tyr Val Asn Leu Ala Pro Ala Val Asn Val Gly Gln His Leu Val Val Asp Leu Ser Thr Gln Ile Phe Cys His Asn Asp Tyr Pro Glu Thr Ile Thr Asp Tyr Val Thr Leu Gln Arg Gly Ser Ala Tyr Gly Gly Val Leu Ser Asn Phe Ser Gly Thr Val Lys Tyr Ser Gly Ser Ser Tyr Pro Phe Pro Thr Thr Ser Glu Thr Leu 105 Arg Val Val Tyr Asn Ser Arg Thr Asp Lys Pro Trp Pro Val Ala Leu Tyr Leu Thr Pro Val Ser Ser Ala Gly Gly Val Ala Ile Lys Ala Gly 135 Ser Leu Ile Ala Val Leu Ile Leu Arg Gln Thr Asn Asn Tyr Asn Ser 150 155 Asp Asp Phe Gln Phe Val Trp Asn Ile Tyr Ala Asn Asn Asp Val Val 170 Val Pro Thr Gly Gly Cys Asp Val Ser Ala Arg Asp Val Thr Val Thr Leu Pro Asp Tyr Pro Gly Ser Val Pro Ile Pro Leu Thr Val Tyr Cys 200 Ala Lys Ser Gln Asn Leu Gly Tyr Tyr Leu Ser Gly Thr His Ala Asp 210 215 Ala Gly Asn Ser Ile Phe Thr Asn Thr Ala Ser Phe Ser Pro Ala Gln Gly Val Gly Val Gln Leu Thr Arg Asn Gly Thr Ile Ile Pro Ala Asn Asn Thr Val Ser Leu Gly Ala Val Gly Thr Ser Ala Val Ser Leu Gly 265 Leu Thr Ala Asn Tyr Ala Arg Thr Gly Gly Gln Val Thr Ala Gly Asn Val Gln Ser Ile Ile Gly Val Thr Phe Val Tyr Gln 295

(2) INFORMATION FOR SEQ ID NO:31:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 296 amino acids

(B) TYPE: amino acid

(C) STRANDEDNESS: unknown

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:

Met Lys Arg Val Ile Thr Leu Phe Ala Val Leu Leu Met Gly Trp Ser 1 5 10 15

Val Asn Ala Trp Ser Phe Ala Cys Lys Thr Ala Asn Gly Thr Ala Ile 20 25 30

Pro Ile Gly Gly Ser Ala Asn Val Tyr Val Asn Leu Ala Pro Ala 35 40 45

Val Asn Val Gly Gln Asn Leu Val Val Asp Leu Ser Thr Gln Ile Phe 50 55 60

Cys His Asn Asp Tyr Pro Glu Thr Ile Thr Asp Tyr Val Thr Leu Gln 65 70 75 80

Arg Gly Ser Ala Tyr Gly Gly Val Leu Ser Asn Phe Ser Gly Thr Val 85 90 95

Lys Tyr Ser Gly Ser Ser Tyr Pro Phe Pro Thr Thr Ser Glu Thr Pro 100 105 110

Arg Val Val Tyr Asn Ser Arg Thr Asp Lys Pro Trp Pro Val Ala Leu 115 120 125

Tyr Leu Thr Pro Val Ser Ser Ala Gly Lys Ala Gly Ser Leu Ile Ala 130 135 140

Val Leu Ile Leu Arg Gln Thr Asn Asn Tyr Asn Ser Asp Asp Phe Gln 145 150 155 160

Phe Val Trp Asn Ile Tyr Ala Asn Asn Asp Val Val Pro Thr Gly
165 170 175

Gly Cys Asp Val Ser Ala Arg Asp Val Thr Val Thr Leu Pro Asp Tyr 180 185 190

Pro Gly Ser Val Pro Ile Pro Leu Thr Val Tyr Cys Ala Lys Ser Gln 195 200 205

Asn Leu Gly Tyr Tyr Leu Ser Gly Thr His Ala Asp Ala Gly Asn Ser 210 215 220

Ile Phe Thr Asn Thr Ala Ser Phe Ser Pro Ala Gln Gly Val Gly Val 225 230 235 240

Gln Leu Thr Arg Asn Gly Thr Ile Ile Pro Ala Asn Asn Thr Val Ser

245 250 255

Leu Gly Ala Val Gly Thr Ser Ala Val Ser Leu Gly Leu Thr Ala Asn 260 265 270

Tyr Ala Arg Thr Gly Gly Gln Val Thr Ala Gly Asn Val Gln Ser Ile 275 280 285

Ile Gly Val Thr Phe Val Tyr Gln 290 295

(2) INFORMATION FOR SEQ ID NO:32:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 300 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:

Met Lys Arg Val Ile Thr Leu Phe Ala Val Leu Leu Met Gly Trp Ser 1 5 10 15

Val Asn Ala Trp Ser Phe Ala Cys Lys Thr Ala Asn Gly Thr Ala Ile 20 25 30

Pro Ile Gly Gly Ser Ala Asn Val Tyr Val Asn Leu Ala Pro Val 35 40 45

Val Asn Val Gly Gln Asn Leu Val Val Asp Leu Ser Thr Gln Ile Phe 50 55 60

Cys His Asn Asp Tyr Pro Glu Thr Ile Thr Asp Tyr Val Thr Arg Gln 65 70 75 80

Arg Gly Ser Ala Tyr Gly Gly Val Leu Ser Asn Phe Ser Gly Thr Val 85 90 95

Lys Tyr Ser Gly Ser Ser Tyr Pro Phe Pro Thr Thr Ser Glu Thr Pro 100 105 110

Arg Val Val Tyr Asn Ser Arg Thr Asp Lys Pro Trp Pro Val Ala Leu 115 120 125

Tyr Leu Thr Pro Val Ser Ser Ala Gly Gly Val Ala Ile Lys Ala Gly
130 135 140

Ser Leu Ile Ala Val Leu Ile Leu Arg Gln Thr Asn Asn Tyr Asn Ser 145 150 155 160

Asp Asp Phe Gln Phe Val Trp Asn Ile Tyr Ala Asn Asn Asp Val Val 165 170 175

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Val Pro Thr Gly Gly Cys Asp Val Ser Ala Arg Asp Val Thr Val Thr 180 185

Leu Pro Asp Tyr Pro Gly Ser Val Pro Ile Pro Leu Thr Val Tyr Cys 200

Ala Lys Ser Gln Asn Leu Gly Tyr Tyr Leu Ser Gly Thr His Ala Asp 215

Ala Gly Asn Ser Ile Phe Thr Asn Thr Ala Ser Phe Ser Pro Ala Gln

Gly Val Gly Val Gln Leu Thr Arg Asn Gly Thr Ile Ile Pro Ala Asn

Asn Thr Val Ser Leu Gly Ala Val Gly Thr Ser Ala Val Ser Leu Gly 260

Leu Thr Ala Asn Tyr Ala Arg Thr Gly Gly Gln Val Thr Ala Gly Asn 280

Val Gln Ser Ile Ile Gly Val Thr Phe Val Tyr Gln 290 295

(2) INFORMATION FOR SEQ ID NO:33:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 300 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:

Met Lys Arg Val Ile Asn Leu Phe Ala Val Leu Leu Met Gly Trp Ser 5

Val Asn Ala Trp Ser Phe Ala Cys Lys Thr Ala Asn Gly Thr Ala Ile

Pro Ile Gly Gly Ser Ala Asn Val Tyr Val Asn Leu Ala Pro Ala

Val Asn Val Gly Gln Asn Leu Val Val Asp Leu Ser Thr Gln Ile Phe 50

Cys His Asn Asp Tyr Pro Glu Thr Ile Thr Asp Tyr Val Thr Leu Gln

Arg Gly Ser Ala Tyr Gly Gly Val Leu Ser Asn Phe Ser Gly Thr Val

Lys Tyr Ser Gly Ser Ser Tyr Pro Phe Pro Thr Thr Ser Glu Thr Pro 100

Arg Val Val Tyr Asn Ser Arg Thr Asp Lys Pro Trp Pro Val Ala Leu
115 120 125

Tyr Leu Thr Pro Val Ser Ser Ala Gly Gly Val Val Ile Lys Ala Gly 130 135 140

Ser Leu Ile Ala Val Leu Ile Leu Arg Gln Thr Asn Asn Tyr Asn Ser 145 150 155 160

Asp Asp Phe Gln Phe Val Trp Asn Ile Tyr Ala Asn Asn Asp Val Val
165 170 175

Val Pro Thr Gly Gly Cys Asp Val Ser Ala Arg Asp Val Thr Val Thr 180 185 190

Leu Pro Asp Tyr Pro Gly Ser Val Pro Ile Pro Leu Thr Val Tyr Cys 195 200 205

Ala Lys Ser Gln Asn Leu Gly Tyr Tyr Leu Ser Gly Thr His Ala Asp 210 215 220

Ala Gly Asn Ser Ile Phe Thr Asn Thr Ala Ser Phe Ser Pro Ala Gln 225 230 235 240

Gly Val Gly Val Gln Leu Thr Arg Asn Gly Thr Ile Ile Pro Ala Asn 245 250 255

Asn Thr Val Ser Leu Gly Ala Val Gly Thr Ser Ala Val Ser Leu Gly 260 265 270

Leu Thr Ala Asn Tyr Ala Arg Thr Gly Gly Gln Val Thr Ala Gly Asn 275 280 285

Val Gln Ser Ile Ile Gly Val Thr Phe Val Tyr Gln 290 295 300

(2) INFORMATION FOR SEQ ID NO:34:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 300 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:

Met Lys Arg Val Ile Thr Leu Phe Ala Val Leu Leu Met Gly Trp Ser

1 10 15

Val Asn Ala Trp Ser Phe Ala Cys Lys Thr Ala Asn Gly Thr Ala Ile 20 25 30

Pro Ile Gly Gly Ser Ala Asn Val Tyr Val Asn Leu Ala Pro Ala 35 40 45

Val Asn Val Gly Gln Asn Leu Val Val Asp Leu Ser Thr Gln Ile Phe 55 Cys His Asn Asp Tyr Pro Glu Thr Ile Thr Asp Tyr Val Thr Leu Gln Arg Gly Ser Ala Tyr Gly Gly Val Leu Ser Ser Phe Ser Gly Thr Val Lys Tyr Asn Gly Ser Ser Tyr Pro Phe Pro Thr Thr Ser Glu Thr Pro 105 Arg Val Val Tyr Asn Ser Arg Thr Asp Lys Pro Trp Pro Val Ala Leu 120 Tyr Leu Thr Pro Val Ser Ser Ala Gly Gly Val Ala Ile Lys Ala Gly 130 135 Ser Leu Ile Ala Val Leu Ile Leu Arg Gln Thr Asn Asn Tyr Asn Ser Asp Asp Phe Gln Phe Val Trp Asn Ile Tyr Ala Asn Asn Asp Val Val Val Pro Thr Gly Gly Cys Asp Val Ser Ala Arg Asp Val Thr Val Thr 180 Leu Pro Asp Tyr Pro Gly Ser Val Pro Ile Pro Leu Thr Val Tyr Cys 200 Ala Lys Ser Gln Asn Leu Gly Tyr Tyr Leu Ser Gly Thr His Ala Asp 210 215 Ala Gly Asn Ser Ile Phe Thr Asn Thr Ala Ser Phe Ser Pro Ala Gln Gly Val Gly Val Gln Leu Thr Arg Asn Gly Thr Ile Ile Pro Ala Asn 250 Asn Thr Val Ser Leu Gly Ala Val Gly Thr Ser Ala Val Ser Leu Gly 260 265 Leu Thr Ala Asn Tyr Ala Arg Thr Gly Gly Gln Val Thr Ala Gly Asn

Val Gln Ser Ile Ile Gly Val Thr Phe Val Tyr Gln 290 295 300

(2) INFORMATION FOR SEQ ID NO:35:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 300 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide

(xi)	SEQUENCE	DESCRIPTION:	SEQ	ID	NO:	: 35	:
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Met Lys Arg Val Ile Thr Leu Phe Ala Val Leu Leu Met Gly Trp Ser 1 5 10 15

Val Asn Ala Trp Ser Phe Ala Cys Lys Thr Ala Asn Gly Thr Ala Ile 20 25 30

Pro Ile Gly Gly Ser Ala Asn Val Tyr Val Asn Leu Ala Pro Ala 35 40 45

Val Asn Val Gly Gln Asn Leu Val Val Asp Leu Ser Thr Gln Ile Phe 50 55 60

Cys His Asn Asp Tyr Pro Glu Thr Ile Thr Asp Tyr Val Thr Leu Gln 65 70 75 80

Arg Gly Ser Ala Tyr Gly Gly Val Leu Ser Asn Phe Ser Gly Thr Val 85 90 95

Lys Tyr Ser Gly Ser Ser Tyr Pro Phe Pro Thr Thr Ser Glu Thr Pro 100 105 110

Arg Val Val Tyr Asn Ser Arg Thr Asp Lys Pro Trp Pro Val Ala Leu 115 120 125

Tyr Leu Thr Pro Val Ser Ser Ala Gly Gly Val Ala Ile Lys Ala Gly
130 135 140

Ser Leu Ile Ala Val Leu Ile Leu Arg Gln Thr Asn Asn Tyr Asn Ser 145 150 155 160

Asp Asp Phe Gln Phe Val Trp Asn Ile Tyr Ala Asn Asn Asp Val Val 165 170 175

Val Pro Thr Gly Gly Cys Asp Val Ser Ala His Asp Val Thr Val Thr 180 185 190

Leu Pro Asp Tyr Pro Gly Ser Val Pro Ile Pro Leu Thr Val Tyr Cys 195 200 205

Ala Lys Ser Gln Asn Leu Gly Tyr Tyr Leu Ser Gly Thr His Ala Asp 210 215 220

Ala Gly Asn Ser Ile Phe Thr Asn Thr Ala Ser Phe Ser Pro Ala Gln 225 230 235 240

Gly Val Gly Val Gln Leu Thr Arg Asn Gly Thr Ile Ile Pro Ala Asn 245 250 255

Asn Thr Val Ser Leu Gly Ala Val Gly Thr Ser Ala Val Ser Leu Gly 260 265 270

Leu Thr Ala Asn Tyr Ala Arg Thr Gly Gly Gln Val Thr Ala Gly Asn 275 280 285

Val Gln Ser Ile Ile Gly Val Thr Phe Val Tyr Gln

295

290

300

- (2) INFORMATION FOR SEQ ID NO:36:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 300 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:

Met Lys Arg Val Ile Thr Leu Phe Ala Val Leu Leu Met Gly Trp Ser 1 5 10 15

Val Asn Ala Trp Ser Phe Ala Cys Lys Thr Ala Asn Gly Thr Ala Ile 20 25 30

Pro Ile Gly Gly Ser Ala Asn Val Tyr Val Asn Leu Ala Pro Ala 35 40 45

Val Asn Val Gly Gln Asn Leu Val Val Asp Leu Ser Thr Gln Ile Phe 50 55 60

Cys His Asn Asp Tyr Pro Glu Thr Ile Thr Asp Tyr Val Thr Leu Gln 65 70 75 80

Arg Gly Ser Ala Tyr Gly Gly Val Leu Ser Asn Phe Ser Gly Thr Val 85 90 95

Lys Tyr Ser Gly Ser Ser Tyr Pro Phe Pro Thr Thr Ser Glu Thr Pro
100 105 110

Arg Val Val Tyr Asn Ser Arg Thr Asp Lys Pro Trp Pro Val Ala Arg 115 120 125

Tyr Leu Thr Pro Val Ser Ser Ala Gly Gly Val Ala Ile Lys Ala Gly
130 135 140

Ser Leu Ile Ala Val Leu Ile Leu Arg Gln Thr Asn Asn Tyr Asn Ser 145 150 155 160

Asp Asp Phe Gln Phe Val Trp Asn Ile Tyr Ala Asn Asn Asp Val Val
165 170 175

Val Pro Thr Gly Gly Cys Asp Val Ser Ala Arg Asp Val Thr Val Thr 180 185 190

Leu Pro Asp Tyr Pro Gly Ser Val Pro Ile Pro Leu Thr Val Tyr Cys
195 200 205

Ala Lys Ser Gln Asn Leu Gly Tyr Tyr Leu Ser Gly Thr His Ala Asp 210 215 220

Ala Gly Asn Ser Ile Phe Thr Asn Thr Ala Ser Phe Ser Pro Ala Gln 225 230 235 240

Gly Val Gly Val Gln Leu Thr Arg Asn Gly Thr Ile Ile Pro Ala Asn 245 250 255

Asn Thr Val Ser Leu Gly Ala Val Gly Thr Ser Ala Val Ser Leu Gly 260 265 270

Leu Thr Ala Asn Tyr Ala Arg Thr Gly Gly Gln Val Thr Ala Gly Asn 275 280 285

Val Gln Ser Ile Ile Gly Val Thr Phe Val Tyr Gln 290 295 300

(2) INFORMATION FOR SEQ ID NO:37:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 300 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:

Met Lys Arg Val Ile Thr Leu Phe Ala Val Leu Leu Met Gly Trp Ser 1 5 10 15

Val Asn Ala Trp Ser Phe Ala Cys Lys Thr Ala Asn Gly Thr Ala Ile 20 25 30

Pro Ile Gly Gly Ser Ala Asn Val Tyr Val Asn Leu Ala Pro Ala
35 40 45

Val Asn Val Gly Gln Asn Leu Val Val Asp Leu Ser Thr Gln Ile Phe 50 55 60

Cys His Asn Asp Tyr Pro Glu Thr Ile Thr Asp Tyr Val Thr Leu Gln 65 70 75 80

Arg Gly Ser Ala Tyr Gly Gly Val Leu Ser Asn Phe Ser Gly Thr Val 85 90 95

Lys Tyr Ser Gly Ser Ser Tyr Pro Phe Pro Thr Thr Ser Glu Thr Pro 100 105 110

Arg Val Val Tyr Asn Ser Arg Thr Asp Lys Pro Trp Pro Val Ala Leu 115 120 125

Tyr Leu Thr Pro Val Ser Ser Ala Gly Gly Val Ala Ile Lys Ala Gly
130 135 140

Ser Leu Ile Ala Val Leu Ile Leu Arg Gln Thr Asn Asn Tyr Asn Ser 145 150 155 160

Asp Asp Phe Gln Phe Val Trp Asn Ile Tyr Ala Asn Asn Asp Val Val
165 170 175

Val Pro Thr Gly Gly Cys Asp Val Ser Ala His Asp Val Thr Val Thr 180 185 190

Leu Pro Asp Tyr Pro Gly Ser Val Pro Ile Pro Leu Thr Val Tyr Cys 195 200 205

Ala Lys Ser Gln Asn Leu Gly Tyr Tyr Leu Ser Gly Thr His Ala Asp 210 215 220

Ala Gly Asn Ser Ile Phe Thr Asn Thr Ala Ser Phe Ser Pro Ala Gln 225 230 235 240

Gly Val Gly Val Gln Leu Thr Arg Asn Gly Thr Ile Ile Pro Ala Asn 245 250 255

Asn Thr Val Ser Leu Gly Ala Val Gly Thr Ser Ala Val Ser Leu Gly 260 265 270

Leu Thr Ala Asn Tyr Ala Arg Thr Gly Gly Gln Val Thr Ala Gly Asn 275 280 285

Val Gln Ser Ile Ile Gly Val Thr Phe Val Tyr Gln 290 295 300

(2) INFORMATION FOR SEQ ID NO:38:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 300 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:

Met Lys Arg Val Ile Thr Leu Phe Ala Val Leu Leu Met Gly Trp Ser

1 10 15

Val Asn Ala Trp Ser Phe Ala Cys Lys Thr Ala Asn Gly Thr Ala Ile 20 25 30

Pro Ile Gly Gly Ser Ala Asn Val Tyr Val Asn Leu Ala Pro Ala 35 40 45

Val Asn Val Gly Gln Asn Leu Val Val Asp Leu Ser Thr Gln Ile Phe 50 55 60

Cys His Asn Asp Tyr Pro Glu Thr Ile Thr Asp Tyr Val Thr Leu Gln 65 70 75 80

Arg Gly Ser Ala Tyr Gly Gly Val Leu Ser Asn Phe Ser Gly Thr Val 85 90 95

PCT/DK95/00042 WO 95/20657 111

Lys Tyr Ser Gly Ser Ser Tyr Pro Phe Pro Thr Thr Ser Glu Thr Pro 110 100 105

Arg Val Val Tyr Asn Ser Arg Thr Asp Lys Pro Trp Pro Val Ala Leu 120

Tyr Leu Thr Pro Val Ser Ser Ala Gly Gly Val Ala Ile Lys Ala Gly

Ser Leu Ile Ala Val Leu Ile Leu Arg Gln Thr Asn Asn Tyr Asn Ser

Asp Asp Phe Gln Phe Val Trp Asn Ile Tyr Ala Asn Asn Asp Val Val 170

Val Pro Thr Gly Gly Cys Asp Val Ser Ala His Asp Val Thr Val Thr

Leu Pro Asp Tyr Pro Gly Ser Val Pro Ile Pro Leu Thr Val Tyr Cys 200

Ala Lys Ser Gln Asn Leu Gly Tyr Tyr Leu Ser Gly Thr His Ala Asp 215 220

Ala Gly Asn Ser Ile Phe Thr Asn Thr Ala Ser Phe Ser Pro Ala Gln 230

Gly Val Gly Val Gln Leu Thr Arg Asn Gly Thr Ile Ile Pro Ala Asn 250

Asn Thr Val Ser Leu Gly Ala Val Gly Thr Ser Ala Val Ser Leu Gly 260 265

Leu Thr Ala Asn Tyr Ala Arg Thr Gly Gly Gln Val Thr Ala Gly Asn 280

Val Gln Ser Ile Ile Gly Val Thr Phe Val Tyr Gln 290 295

(2) INFORMATION FOR SEQ ID NO:39:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 300 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:

Met Lys Arg Val Ile Thr Leu Phe Ala Val Leu Leu Met Gly Trp Ser

Val Asn Ala Trp Ser Phe Ala Cys Lys Thr Ala Asn Gly Thr Ala Ile 25

Pro Ile Gly Gly Ser Ala Asn Val Tyr Val Asn Leu Ala Pro Ala 35 40 45

Val Asn Val Gly Gln Asn Leu Val Val Asp Leu Ser Thr Gln Ile Phe 50 55 60

Cys His Asn Asp Tyr Pro Glu Thr Ile Thr Asp Tyr Val Thr Leu Gln 65 70 75 80

Arg Gly Ser Ala Tyr Gly Gly Val Leu Ser Ser Phe Ser Gly Thr Val 85 90 95

Lys Tyr Asn Gly Ser Ser Tyr Pro Phe Pro Thr Thr Ser Glu Thr Pro 100 105 110

Arg Val Val Tyr Asn Ser Arg Thr Asp Lys Pro Trp Pro Val Ala Leu 115 120 125

Tyr Leu Thr Pro Val Ser Ser Ala Gly Gly Val Ala Ile Lys Ala Gly
130 135 140

Ser Leu Ile Ala Val Leu Ile Leu Arg Gln Thr Asn Asn Tyr Asn Ser I45 150 155 160

Asp Asp Phe Gln Phe Val Trp Asn Ile Tyr Ala Asn Asn Asp Val Val 165 170 175

Val Pro Thr Gly Gly Cys Asp Ala Ser Ala Arg Asp Val Thr Val Thr 180 185 190

Leu Pro Asp Tyr Pro Gly Ser Val Pro Ile Pro Leu Thr Val Tyr Cys
195 200 205

Ala Lys Ser Gln Asn Leu Gly Tyr Tyr Leu Ser Gly Thr His Ala Asp 210 215 220

Ala Gly Asn Ser Ile Phe Thr Asn Thr Ala Ser Phe Ser Pro Ala Gln 225 230 235 240

Gly Val Gly Val Gln Leu Thr Arg Asn Gly Thr Ile Ile Pro Ala Asn 245 250 255

Asn Thr Val Ser Leu Gly Ala Val Gly Thr Ser Ala Val Ser Leu Gly
260 265 270

Leu Thr Ala Asn Tyr Ala Arg Thr Gly Gly Gln Val Thr Ala Gly Asn 275 280 285

Val Gln Ser Ile Ile Gly Val Thr Phe Val Tyr Gln 290 295 300

(2) INFORMATION FOR SEQ ID NO:40:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 300 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi)	SEQ	JENC	E DE	SCRI	PTIO	N: S1	EQ II	ON C	:40:						
Met 1	Lys	Arg	Val	Ile 5	Thr	Leu	Phe	Ala	Val 10	Leu	Leu	Met	Gly	Trp 15	Ser
Val	Asn	Ala	Trp 20	Ser	Phe	Ala	Cys	Lys 25	Thr	Ala	Asn	Gly	Thr 30	Ala	Ile
Pro	Ile	Gly 35	Gly	Gly	Ser	Ala	Asn 40	Val	Tyr	Val	Asn	Leu 45	Ala	Pro	Ala
Val	Asn 50	Val	Gly	Gln	Asn	Leu 55	Val	Val	Asp	Leu	Ser 60	Thr	Gln	Ile	Phe
Сув 65	His	Asn	Asp	Tyr	Pro 70	Glu	Thr	Ile	Thr	Авр 75	Tyr	Val	Thr	Leu	Gln 80
Arg	Gly	Ser	Ala	Tyr 85	Gly	Asp	Val	Leu	Ser 90	Ser	Phe	Ser	Gly	Thr 95	Val
Lys	Tyr	Asn	Gly 100	Ser	Ser	Tyr	Pro	Phe 105	Pro	Thr	Thr	Ser	Glu 110	Thr	Pro
Arg	Val	Val 115	Tyr	Asn	Ser	Arg	Thr 120	qaA	Lys	Pro	Trp	Pro 125	Val	Ala	Leu
Tyr	Leu 130	Thr	Pro	Val	Ser	Ser 135	Ala	Gly	Gly	Val	Ala 140	Ile	Lys	Ala	Gly
Ser 145	Leu	Ile	Ala	Val	Leu 150	Ile	Leu	Arg	Gln	Thr 155	Asn	Asn	Tyr	Asn	Ser 160
qaA	Asp	Phe _.	Gln	Phe 165	Val	Trp	Asn	Ile	Tyr 170	Ala	Asn	Asn	qaA	Val 175	Val
Val	Pro	Thr	Gly 180	Gly	Сув	Asp	Ala	Ser 185	Ala	Arg	Asp	Val	Thr 190	Val	Thr
Leu	Pro	Asp 195	Tyr	Pro	Gly	Ser	Val 200	Pro	Ile	Pro	Leu	Thr 205	Val	Tyr	Сув
Ala	Lys 210	Ser	Gln	Asn	Leu	Gly 215	Tyr	Tyr	Leu	Ser	Gly 220	Thr	His	Ala	Asp
Ala 225	Gly	Asn	Ser	Ile	Phe 230	Thr	Asn	Thr	Ala	Ser 235	Phe	Ser	Pro	Ala	Gln 240
Gly	Val	Gly	Val	Gln 245	Leu	Thr	Arg	Asn	Gly 250	Thr	Ile	Ile	Pro	Ala 255	Asn
Asn	Thr	Val	Ser 260	Leu	Gly	Ala	Val	Gly 265	Thr	Ser	Ala	Val	Ser 270	Leu	Gly
Leu	Thr	Ala	Asn	Tyr	Ala	Arg	Thr	Gly	Gly	Gln	Val	Thr	Ala	Gly	Asn

280

114

Val Gln Ser Ile Ile Gly Val Thr Phe Val Tyr Gln

290 295

(2) INFORMATION FOR SEQ ID NO:41:

275

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 900 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:

IGCTGTA CTGCTGATGG GCT	GGTCGGT AAATGCCTGG 60
rggtacc gctatcccta tigo	GCGGTGG CAGCGCCAAT 120
CGTGAAT GTGGGGCAAA ACC	TGGTCGT GGATCTTTCG 180
TTATCCG GAAACCATTA CAG	ACTATGT CACACTGCAA 240
GTTATCT AATTTTTCCG GGA	CCGTAAA ATATAGTGGC 300
CAGCGAA ACGCCGCGCG TTG	TITATAA TTCGAGAACG 360
FTATTTG ACGCCTGTGA GCAC	GTGCGGG CGGGGTGGCG 420
CGTGCTT ATTTTGCGAC AGA	CCAACAA CTATAACAGC 480
FATTTAC GCCAATAATG ATG	TGGTGGT GCCTACTGGC 540
TGTCACC GTTACTCTGC CGG	ACTACCG TGGTTCAGTG 600
rgcgaaa agccaaaacc tgg	GGTATTA CCTCTCCGGC 660
SATTITC ACCAATACCG CGT	CGTTTTC ACCTGCACAG 720
CAACGGT ACGATTATTC CAG	CGAATAA CACGGTATCG 780
egtgagt ctgggattaa cgg	CAAATTA TGCACGTACC 840
GTGCAA TCGATTATTG GCGT	TGACTIT TGTTTATCAA 900

(2) INFORMATION FOR SEQ ID NO:42:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 900 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:42: ATGAAACGAG TTATTACCCT GTTTGCTGTA CTGCTGATGG GCTGGTCGGT AAATGCCTGG 60 TCATTCGCCT GTAAAACCGC CAATGGTACC GCAATCCCTA TTGGCGGTGG CAGCGCCAAT 120 GTTTATGTAA ACCTTGCGCC TGCCGTGAAT GTGGGGCAAA ACCTGGTCGT AGATCTTTCG 180 ACGCAAATCT TTTGCCATAA CGATTACCCA GAAACCATTA CAGACTATGT CACACTGCAA 240 CGAGGTTCGG CTTATGGCGG CGTGTTATCT AGTTTTTCCG GGACCGTAAA ATATAATGGC 300 AGTAGCTATC CTTTCCCTAC TACCAGCGAA ACGCCGCGGG TTGTTTATAA TTCGAGAACG 360 GATAAGCCGT GGCCGGTGGC GCTTTATTTG ACGCCGGTGA GCAGTGCGGG GGGAGTGGCG 420 ATTAAAGCTG GCTCATTAAT TGCCGTGCTT ATTTTGCGAC AGACCAACAA CTATAACAGC 480 GATGATTTCC AGTITGTGTG GAATATTTAC GCCAATAATG ATGTGGTGGT GCCCACTGGC 540 GGCTGCGATG TTTCTGCTCG TGATGTCACC GTTACTCTGC CGGACTACCC TGGTTCAGTG 600 CCGATTCCTC TTACCGTTTA TTGTGCGAAA AGCCAAAACC TGGGGTATTA CCTCTCCGGC 660 ACACACGCAG ATGCGGGCAA CTCGATTTTC ACCAATACCG CGTCGTTTTC ACCCGCGCAG 720 GGCGTCGGCG TACAGTTGAC GCGCAACGGT ACGATTATTC CAGCGAATAA CACGGTATCG 780 TTAGGAGCAG TAGGGACTTC GGCGGTAAGT CTGGGATTAA CGGCAAATTA CGCACGTACC 840 GGAGGGCAGG TGACTGCAGG GAATGTGCAA TCGATTATTG GCGTGACTTT TGTTTATCAA 900

(2) INFORMATION FOR SEQ ID NO:43:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 900 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:43:

ATGAAACGAG TTATTACCCT GTTTGCTGTA CTGCTGATGG GCTGGTCGGT AAATGCCTGG 60

TCATTCGCCT GTAAAACCGC CAATGGTACC GCAATCCCTA TTGGCGGTGG CAGCGCCAAT 120

GTTTATGTAA ACCTTGCGCC TGCCGTGAAT GTGGGGCAAA ACCTGGTCGT AGATCTTTCG 180

ACGCAAATCT TTTGCCATAA CGATTACCCA GAAACCATTA CAGACTATGT CACACTGCAA 240



CGAGGTTCGG CTTATGGCGG CGTGTTATCT AGTTTTTCCG GGACCGTAAA ATATAATGGC 300 AGTAGCTATC CTTTCCCTAC TACCAGCGAA ACGCCGCGGG TTGTTTATAA TTCGAGAACG 360 GATAAGCCGT GGCCGGTGGC GCTTTATTTG ACGCCGGTGA GCAGTGCGGG GGGAGTGGCG 420 ATTAAAGCTG GCTCATTAAT TGCCGTGCTT ATTTTGCGAC AGACCAACAA CTATAACAGC 480 GATGATTTCC AGTTTGTGTG GAATATTTAC GCCAATAATG ATGTGGTGGT GCCCACTGGC 540 GGCTGTGATG CTTCTGCTCG TGATGTCACC GTTACTCTGC CGGACTACCC TGGTTCAGTG 600 CCGATTCCTC TTACCGTTTA TTGTGCGAAA AGCCAAAACC TGGGGTATTA CCTATCCGGC 660 ACACATGCAG ATGCGGGCAA CTCGATTTTC ACCAATACCG CGTCGTTTTC ACCCGCGCAG 720 GGCGTCGGCG TACAGTTGAC GCGCAACGGT ACGATTATTC CAGCGAATAA CACGGTATCG 780 TTAGGAGCAG TAGGGACTTC GGCGGTGAGT CTGGGATTAA CGGCAAATTA TGCACGTACC 840 GGAGGGCAGG TGACTGCAGG GAATGTGCAA TCGATTATTG GCGTGACTTT TGTTTATCAA 900

(2) INFORMATION FOR SEQ ID NO:44:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 900 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:44:

ATGAAACGAG	TTATTACCCT	GTTTGCTGTA	CTGCTGATGG	GCTGGTCGGT	AAATGCCTGG	60
TCATTCGCCT	GTAAAACCGC	CAATGGTACC	GCTATCCCTA	TTGGCGGTGG	CAGCGCCAAT	120
GTTTATGTAA	ACCTTGCGCC	TGCCGTGAAT	GTGGGGCAAA	ACCTGGTCGT	GGATCTTTCG	180
ACGCAAATCT	TTTGCCATAA	CGATTACCCG	GAAACCATTA	CAGACTATGT	CACACTGCAA	240
CGAGGTTCGG	CTTATGGCGG	CGTGTTATCT	AGTTTTTCCG	AGACCGTAAA	ATATAATGGC	300
AGTAGCTATC	CTTTCCCTAC	TACCAGCGAA	ACGCCGCGGG	TTGTTTATAA	TTCGAGAACG	360
GATAAGCCGT	GGCCGGTGGC	GCTTTATTTG	ACGCCTGTGA	GCAGTGCGGG	GGGAGTGGCG	420
ATTAAAGCTG	GCTCATTAAT	TGCCGTGCTT	ATTTTGCGAC	AGACCAACAA	CTATAACAGC	480
GATGATTTCC	AGTTTGTGTG	GAATATTTAC	GCCAATAATG	ATGTGGTGGT	GCCCACTGGC	540
GGCTGTGATG	TTTCTGCTCG	TGATGTCACC	GTTACTTTGC	CGGACTACCC	TGGTTCAGTG	600
CCGATTCCTC	TTACCGTTTA	TTGTGCGAAA	AGCCAAAACC	TGGGGTATTA	CCTCTCCGGC	660

ACAGACGCAG	ATGCGGGCAA	CTCGATTTTC	ACCAATACCG	CGTCGTTTTC	ACCTGCACAG	720
GGCGTCGGCG	TACAGTTGAC	GCGCAACGGT	ACGATTATTC	CAGCGAATAA	CACGGTATCG	780
TTAGGAGCAG	TAGGGACTTC	GGCGGTAAGT	CTGGGATTAA	CGGCAAATTA	CGCACGTACC	840
GGAGGGCAGG	TGACTGCAGG	GAATGTGCAA	TCGATTATTG	GCGTGACTTT	TGTTTATCAA	900

(2) INFORMATION FOR SEQ ID NO:45:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 900 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:45:

ATGAAACGAG	TTATTACCCT	GTTTGCTGTA	CTGCTGATGG	GCTGGTCGGT	AAATGCCTGG	60
TCATTCGCCT	GTAAAACCGC	CAATGGTACC	GCAATCCCTA	TTGGCGGTGG	CAGCGCCAAT	120
GTTTATGTAA	ACCTTGCGCC	TGCCGTGAAT	GTGGGGCAAA	ACCTGGTCGT	AGATCTTTCG	180
ACGCAAATCT	TTTGCCATAA	CGATTACCCA	GAAACCATTA	CAGACTATGT	CACACTGCAA	240
CGAGGTTCGG	CTTATGGCGA	CGTGTTATCT	AGTTTTTCCG	GGACCGTAAA	ATATAATGGC	300
AGTAGCTATC	CTTTCCCTAC	TACCAGCGAA	ACGCCGCGGG	TTGTTTATAA	TTCGAGAACG	360
GATAAGCCGT	GGCCGGTGGC	GCTTTATTTG	ACGCCGGTGA	GCAGTGCGGG	GGGAGTGGCG	420
ATTAAAGCTG	GCTCATTAAT	TGCCGTGCTT	ATTTTGCGAC	AGACCAACAA	CTATAACAGC	480
GATGATTTCC	AGTTTGTGTG	GAATATTTAC	GCCAATAATG	ATGTGGTGGT	GCCCACTGGC	540
GGCTGTGATG	TCTCTGCTCG	TGATGTCACC	GTTACTCTGC	CGGACTACCC	TGGTTCAGTG	600
CCGATTCCTC	TTACCGTTTA	TTGTGCGAAA	AGCCAAAACC	TGGGGTATTA	CCTATCCGGC	660
ACACACGCAG	ATGCGGGCAA	CTCGATTTTC	ACCAATACCG	CGTCGTTTTC	ACCCGCGCAG	720
GGCGTCGGCG	TACAGTTGAC	GCGCAACGGT	ACGATTATTC	CAGCGAATAA	CACGGTATCG	780
TTAGGAGCAG	TAGGGACTTC	GGCGGTAAGT	CTGGGATTAA	CGGCAAATTA	CGCACGTACC	840
GGAGGGCAGG	TGACCGCAGG	GAATGTGCAA	TCGATTATTG	GCGTGACTTT	TGTTTATCAA	900

(2) INFORMATION FOR SEQ ID NO:46:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 900 base pairs

(B)	TYPE: nucleic	acid
(C)	STRANDEDNESS:	single
	TODOLOGY. 14-	

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:46:

ATGAAACGAG	TTATTACCCT	GTTTGCTGTA	CTGCTGATGG	GCTGGTCGGT	AAATGCCTGG	60
TCATTCGCCT	GTAAAACCGC	CAATGGTACC	GCTATTCCTA	TTGGCGGTGG	CAGCGCTAAT	120
GTTTATGTAA	ACCTTGCGCC	TGCCGTGAAT	GTGGGGCAAA	ACCTGGTCGT	AGATCTTTCG	180
ACGCAAATCT	TTTGCCATAA	CGATTATCCG	GAAACCATTA	CAGACTATGT	CACACTGCAA	240
CGAGGCTCGG	CTTATGGCGG	CGTGTTATCT	AATTTTCCG	GGACCGTAAA	ATATAGTGGC	300
AGTAGCTATC	CATTCCCGAC	TACCAGCGAA	ACGCCGCGGG	TIGTITATAA	TTCGAGAACG	360
GATAAGCCGT	GGCCGGTGGC	GCTTTATTTG	ACGCCTGTGA	GCAGTGCGGG	TGGGGTGGCG	420
ATTAAAGCTG	GCTCATTAAT	TGCCGTGCTT	ATTTTGCGAC	AGACCAACAA	CTATAACAGC	480
GATGATTTCC	AGTTTGTGTG	GAATATTTAC	GCCAATAATG	ATGTGGTGGT	GCCTACTGGC	540
GGCTGCGATG	TTTCTGCTCA	TGATGTCACC	GTTACTCTGC	CGGACTACCC	TGGTTCAGTG	600
CCAATTCCTC	TTACCGTTTA	TTGTGCGAAA	AGCCAAAACC	TGGGGTATTA	CCTCTCCGGC	660
ACACACGCAG	ATGCGGGCAA	CTCGATTTTC	ACCAATACCG	CGTCGTTTTC	ACCAGCGCAG	720
GGCGTCGGCG	TACAGTTGAC	GCGCAACGGT	ACGATTATTC	CAGCGAATAA	CACGGTATCG	780
TTAGGAGCAG	TAGGGACTTC	GGCGGTAAGT	CTGGGATTAA	CGGCAAATTA	CGCACGTACC	840
GGAGGGCAGG	TGACTGCAGG	GAATGTGCAA	TCGATTATTG	GCGTGACTTT	TGTTTATCAA	900

(2) INFORMATION FOR SEQ ID NO:47:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 900 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:

ATGAAACGAG TTATTACCCT GTTTGCTGTA CTGCTGATGG GCTGGTCGGT AAATGCCTGG 60 TCATTCGCCT GTAAAACCGC CAATGGTACC GCTATTCCTA TTGGCGGTGG CAGCGCTAAT 120 WO 95/20657 PCT/DK95/00042 119

GTTTATGTAA	ACCTTGCGCC	TGCCGTGAAT	GTGGGGCAAA	ACCTGGTCGT	AGATCTTTCG	180
ACGCAAATCT	TTTGCCATAA	CGATTATCCG	GAAACCATTA	CAGACTATGT	CACACTGCAA	240
CGAGGCTCGG	CTTATGGCGG	CGTGTTATCT	AATTTTTCCG	GGACCGTAAA	ATATAGTGGC	300
AGTAGCTATC	CATTTCCGAC	TACCAGCGAA	ACGCCGCGG	TTGTTTATAA	TTCGAGAACG	360
GATAAGCCGT	GGCCGGTGGC	GCTTTATTTG	ACGCCTGTGA	GCAGTGCGGG	TGGGGTGGCG	420
ATTAAAGCTG	GCTCATTAAT	TGCCGTGCTT	ATTTTGCGAC	AGACCAACAA	CTATAACAGC	480
GATGATTTCC	AGTTTGTGTG	GAATATTTAC	GCCAATAATG	ATGTGGTGGT	GCCTACTGGC	540
GGCTGCGATG	TTTCTGCTCA	TGATGTCACC	GTTACTCTGC	CGGACTACCC	TGGTTCAGTG	600
CCAATTCCTC	TTACCGTTTA	TTGTGCGAAA	AGCCAAAACC	TGGGGTATTA	CCTCTCCGGC	660
ACACACGCAG	ATGCGGGCAA	CTCGATTTTC	ACCAATACCG	CGTCGTTTTC	ACCAGCGCAG	720
GGCGTCGGCG	TACAGTTGAC	GCGCAACGGT	ACGATTATTC.	CAGCGAATAA	CACGGTATCG	780
TTAGGAGCAG	TAGGGACTTC	GGCGGTAAGT	CTGGGATTAA	CGGCAAATTA	CGCACGTACC	84 Õ
GGAGGGCAGG	TGACTGCAGG	GAATGTGCAA	TCGATTATTG	GCGTGACTTT	TGTTTATCAA	900

(2) INFORMATION FOR SEQ ID NO:48:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 888 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:48:

ATGAAACGAG	TTATTACCCT	GTTTGCTGTA	CTGCTGATGG	GCTGGTCGGT	AAATGCCTGG	60
TCATTCGCCT	GTAAAACCGC	CAATGGTACC	GCTATCCCTA	TTGGCGGTGG	CAGCGCCAAT	120
GTTTATGTAA	ACCTTGCGCC	CGCCGTGAAT	GTGGGGCAAA	ACCTGGTCGT	GGATCTTTCG	180
ACGCAAATCT	TTTGCCATAA	CGATTATCCG	GAAACCATTA	CAGACTATGT	CACACTGCAA	240
CGAGGCTCGG	CTTATGGCGG	CGTGTTATCT	AATTTTTCCG	GGACCGTAAA	ATATAGTGGC	300
AGTAGCTATC	CATTTCCTAC	CACCAGCGAA	ACGCCGCGCG	TTGTTTATAA	TTCGAGAACG	360
GATAAGCCGT	GGCCGGTGGC	GCTTTATTTG	ACGCCTGTGA	GCAGTGCGGG	TAAAGCTGGC	420
TCATTAATTG	CCGTGCTTAT	TTTGCGACAG	ACCAACAACT	ATAACAGCGA	TGATTTCCAG	480
TTTGTGTGGA	ATATTTACGC	CAATAATGAT	GTGGTGGTGC	CTACTGGCGG	CTGCGATGTT	540

TCTGCTCG	GG	ATGTCACCGT	TACTCTGCCG	GACTACCCTG	GTTCAGTGCC	AATTCCTCTT	60 0
ACCGTTTA	TT	GTGCGAAAAG	CCAAAACCTG	GGGTATTACC	TCTCCGGCAC	ACACGCAGAT	660
GCGGGCAA	CT	CGATTTTCAC	CAATACCGCG	TCGTTTTCAC	CTGCACAGGG	CGTCGGCGTA	720
CAGTTGAC	GC	GCAACGGTAC	GATTATTCCA	GCGAATAACA	CGGTATCGTT	AGGAGCAGTA	780
GGGACTTC	:GG	CGGTGAGTTT	GGGATTAACG	GCAAATTATG	CACGTACCGG	AGGGCAGGTG	840
ACTGCAGG	GA.	ATGTGCAATC	GATTATTGGC	GTGACTTTTG	TTTATCAA	•	888

(2) INFORMATION FOR SEQ ID NO:49:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 900 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:49:

ATGAAACGAG	TTATTACCCT	GTTTGCTGTA	CTGCTGATGG	GCTGGTCGGT	AAATGCCTGG	60
TCATTCGCCT	GTAAAACCGC	CAATGGTACC	GCTATCCCTA	TTGGCGGTGG	CAGCGCCAAT	120
GTTTATGTAA	ACCTTGCGCC	CGTCGTGAAT	GTGGGGCAAA	ACCTGGTCGT	GGATCTTTCG	180
ACGCAAATCT	TTTGCCATAA	CGATTATCCG	GAAACCATTA	CAGACTATGT	CACACTGCAA	240
CGAGGCTCGG	CTTATGGCGG	CGTGTTATCT	AATTTTTCCG	GGACCGTAAA	ATATAGTGGC	300
AGTAGCTATC	CATTTCCTAC	CACCAGCGAA	ACGCCGCGCG	TTGTTTATAA	TTCGAGAACG	360
GATAAGCCGT	GGCCGGTGGC	GCTTTATTTG	ACGCCTGTGA	GCAGTGCGGG	CGGGGTGGCG	420
ATTAAAGCTG	GCTCATTAAT	TGCCGTGCTT	ATTTTGCGAC	AGACCAACAA	CTATAACAGC	480
GATGATTTCC	AGTTTGTGTG	GAATATTTAC	GCCAATAATG	ATGTGGTGGT	GCCTACTGGC	540
GGCTGCGATG	TTTCTGCTCG	TGATGTCACC	GTTACTCTGC	CGGACTACCC	TGGTTCAGTG	600
CCAATTCCTC	TTACCGTTTA	TTGTGCGAAA	AGCCAAAACC	TGGGGTATTA	CCTCTCCGGC	660
ACACACGCAG	ATGCGGGCAA	CTCGATTTTC	ACCAATACCG	CGTCGTTTTC	ACCTGCACAG	720
GGCGTCGGCG	TACAGTTGAC	GCGCAACGGT	ACGAATATTC	CAGCGAATAA	CACGGTATCG	780
TTAGGAGCAG	TAGGGACTTC	GGCGGTGAGT	CTGGGATTAA	CGGCAAATTA	TGCACGTACC	840
GGAGGGCAGG	TGACTGCAGG	GAATGTGCAA	TCGATTATTG	GCGTGACTTT	TGTTTATCAA	900

(2) INFORMATION FOR SEQ ID NO:50:

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(i)	SEQUENCE	CHARACTERISTICS	
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- (A) LENGTH: 900 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:50:

ATGAAACGAG	TTATTAACCT	GTTTGCTGTA	CTGCTGATGG	GCTGGTCGGT	AAATGCCTGG	60
TCATTCGCCT	GTAAAACCGC	CAATGGCACC	GCTATCCCTA	TTGGCGGTGG	CAGCGCCAAT	120
GTTTATGTAA	ACCITGCGCC	CGCCGTGAAT	GTGGGGCAAA	ACCTGGTCGT	GGATCTTTCG	180
ACGCAAATCT	TTTGCCATAA	CGATTACCCG	GAAACCATTA	CAGATTATGT	CACACTGCAA	240
CGAGGCTCGG	CTTATGGCGG	CGTGTTATCT	AATTTTTCCG	GGACCGTAAA	ATATAGTGGC	300
AGTAGCTATC	CATTTCCGAC	CACCAGTGAA	ACGCCGCGGG	TIGTITATAA	TTCGAGAACG	360
GATAAGCCGT	GGCCGGTGGC	GCTTTATTTG	ACGCCTGTGA	GCAGTGCGGG	CGGGGTGGTG	420
ATTAAAGCTG	GCTCATTAAT	TGCCGTGCTT	ATTTTGCGAC	AGACCAACAA	CTATAACAGC	480
GATGATTTCC	AGTTTGTGTG	GAATATTTAC	GCCAATAATG	ATGTGGTGGT	GCCCACTGGC	540
GGCTGCGATG	TITCIGCTCG	TGATGTCACC	GTTACTCTGC	CGGACTACCC	TGGTTCAGTG	600
CCGATTCCTC	TTACCGTTTA	TTGTGCGAAA	AGCCAAAACC	TGGGGTATTA	CCTCTCCGGC	660
ACACACGCAG	ATGCGGGCAA	CTCGATTTTC	ACCAATACCG	CGTCGTTTTC	ACCTGCACAG	720
GGCGTCGGCG	TACAGTTGAC	GCGCAACGGT	ACGATTATTC	CAGCGAATAA	CACGGTATCG	780
TTAGGAGCAG	TAGGGACTTC	GGCGGTAAGT	CTGGGATTAA	CGGCAAATTA	CGCACGTACC	840
GGAGGGCAGG	TGACTGCAGG	GAATGTGCAA	TCGATTATTG	CCGTGACTTT	TGTTTATCAA	900

(2) INFORMATION FOR SEQ ID NO:51:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 900 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:51:

ATGAAACGAG TTATTAACCT GTTTGCTGTA CTGCTGATGG GCTGGTCGGT AAATGCCTGG

TCATTCGCCT	GTAAAACCGC	CAATGGCACC	GCTATCCCTA	TTGGCGGTGG	CAGCGCCAAT	120
GTTTATGTAA	ACCTTGCGCC	CGCCGTGAAT	GTGGGGCAAC	ACCTGGTCGT	AGATCTTTCG	180
ACGCAAATCT	TTTGCCATAA	CGATTACCCG	GAAACCATTA	CAGACTATGT	CACACTGCAA	240
CGAGGTTCGG	CTTATGGCGG	CGTGTTATCT	AATTTTTCCG	GGACCGTAAA	ATATAGTGGC	300
AGTAGCTATC	CATTTCCTAC	CACCAGCGAA	ACGCTGCGGG	TTGTTTATAA	TTCGAGAACG	360
GATAAGCCGT	GGCCGGTGGC	GCTTTATTTG	ACGCCTGTGA	GCAGTGCGGG	CGGGGTGGCG	420
ATTAAAGCTG	GCTCATTAAT	TGCCGTGCTT	ATTTTGCGAC	AGACCAACAA	CTATAACAGC	480
GATGATTTCC	AGTTTGTGTG	GAATATTTAC	GCCAATAATG	ATGTGGTGGT	GCCTACTGGC	540
GGCTGCGATG	TTTCTGCTCG	TGATGTCACC	GTTACTCTGC	CGGACTACCC	TGGTTCAGTG	600
CCAATTCCTC	TTACCGTTTA	TTGTGCGAAA	AGCCAAAACC	TGGGGTATTA	CCTCTCCGGC	660
ACACACGCAG	ATGCGGGCAA	CTCGATTTTC	ACCAATACCG	CCTCGTTTTC	ACCAGCGCAG	720
GCGTĊGGCG	TACAGTTGAC	GCGCAACGGT	ACGATTATTC	CAGCGAATAA	CACGGTATCG	780
PTAGGAGCAG	TAGGGACTTC	GGCGGTAAGT	CTGGGATTAA	CGGCAAATTA	CGCACGTACC	840
GGAGGGCAGG	TGACTGCAGG	GAATGTGCAA	TCGATTATTG	GCGTGACTTT	TGTTTATCAA	900

(2) INFORMATION FOR SEQ ID NO:52:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 900 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:

ATGAAACGAG	TTATTACCCT	GTTTGCTGTA	CTGCTGATGG	GCTGGTCGGT	AAATGCCTGG	60
TCATTCGCCT	GTAAAACCGC	CAATGGTACC	GCTATCCCTA	TTGGCGGTGG	CAGCGCTAAT	120
GTTTATGTAA	ACCTTGCGCC	TGCCGTGAAT	GTGGGGCAAA	ACCTGGTCGT	AGATCTITCG	180
ACGCAAATCT	TTTGCCATAA	CGATTATCCG	GAAACCATTA	CAGACTATGT	CACACTGCAA	240
CGAGGCTCGG	CTTATGGCGG	CGTGTTATCT	AATTTTTCCG	GGACCGTAAA	ATATAGTGGC	300
AGTAGCTATC	CATTTCCGAC	TACCAGCGAA	ACGCCGCGG	TTGTTTATAA	TTCGAGAACG	360
GATAAGCCGT	GGCCGGTGGC	GCTTTATTTG	ACGCCTGTGA	GCAGTGCGGG	TGGGGTGGCG	420
ATTAAAGCTG	GCTCATTAAT	TGCCGTGCTT	ATTTTGCGAC	AGACCAACAA	CTATAACAGC	480

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GATGATTTCC	AGTTTGTGTG	GAATATTTAC	GCCAATAATG	ATGTGGTGGT	GCCTACTGGC	540
GGCTGCGATG	TTTCTGCTCA	TGATGTCACC	GTTACTCTGC	CGGACTACCC	TGGTTCAGTG	600
CCAATTCCTC	TTACCGTTTA	TTGTGCGAAA	AGCCAAAACC	TGGGGTATTA	CCTCTCCGGC	660
ACACACGCAG	ATGCGGGCAA	CTCGATTTTC	ACCAATACCG	CGTCGTTTTC	ACCAGCGCAG	720
GGCGTCGGCG	TACAGTTGAC	GCGCAACGGT	ACGATTATTC	CAGCGAATAA	CACGGTATCG	780
TTAGGAGCAG	TAGGGACTTC	GGCGGTAAGT	CTGGGATTAA	CGGCAAATTA	CGCACGTACC	840
GGAGGGCAGG	TGACTGCAGG	GAATGTGCAA	TCGATTATTG	GCGTGACTTT	TGTTTATCAA	900

(2) INFORMATION FOR SEQ ID NO:53:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 900 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:

ATGAAACGAG	TTATTACCCT	GTTTGCTGTA	CTGCTGATGG	GCTGGTCGGT	AAATGCCTGG	60
TCATTCGCCT	GTAAAACCGC	CAATGGTACC	GCTATCCCTA	TTGGCGGTGG	CAGCGCCAAT	120
GTTTATGTAA	ACCTTGCGCC	CGCCGTGAAT	GTGGGGCAAA	ACCTGGTCGT	GGATCTTTCG	180
ACGCAAATCT	TTTGCCATAA	CGATTATCCG	GAAACCATTA	CAGACTATGT	CACACTGCAA	240
CGAGGCTCGG	CTTATGGCGG	CGTGTTATCT	AATTTTTCCG	GGACCGTAAA	ATATAGTGGC	300
AGTAGCTATC	CATTTCCTAC	CACCAGCGAA	ACGCCGCGCG	TTGTTTATAA	TTCGAGAACG	360
GATAAGCCGT	GGCCGGTGGC	GCTTTATTTG	ACGCCTGTGA	GCAGTGCGGG	CGGGGTGGCG	420
ATTAAAGCTG	GCTCATTAAT	TGCCGTGCTT	ATTTTGCGAC	AGACCAACAA	CTATAACAGC	480
GATGATTTCC	AGTTTGTGTG	GAATATTTAC	GCCAATAATG	ATGTGGTGGT	GCCTACTGGC	540
GGCTGCGATG	TTTCTGCTCG	TGATGTCACC	GTTACTCTGC	CGGACTACCC	AGGTTCAGTG	600
CCAATTCCTC	TTACCGTTTA	TTGTGCGAAA	AGCCAAAACC	TGGGGTATTA	CCTCTCCGGC	660
ACACACGCAG	ATGCGGGCAA	CTCGATTTTC	ACCAATACCG	CGTCGTTTTC	ACCTGCACAG	720
GGCGTCGGCG	TACAGTTGAC	GCGCAACGGT	ACGATTATTC	CAGCGAATAA	CACGGTATCG	780
TTAGGAGCAG	TAGGGACTTC	· ·	CTGGGATTAA	CGGCAAATTA	TGCACGTACC	840
GGAGGGCAGG	TGACTGCAGG	GAATGTGCAA	TCGATTATTG	GCGTGACTTT	TGTTTATCAA	900

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(2) INFORMATION FOR SEQ ID NO:54:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 900 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:54:

ATGAAACGAG TTATTACCCT	GTTTGCTGTA	CTGCTGATGG	GCTGGTCGGT	AAATGCCTGG	60
TCATTCGCCT GTAAAACCGC	CAATGGTACC	GCTATCCCTA	TTGGCGGTGG	CAGCGCCAAT	120
GTTTATGTAA ACCTTGCGCC	TGCCGTGAAT	GTGGGGCAAA	ACCTGGTCGT	GGATCTTTCG	180
ACGCAAATCT TITGCCATAA	CGATTACCCG	GAAACCATTA	CAGACTATGT	CACACTGCAA	240
CGAGGTTCGG CTTATGGCGG	CGTGTTATCT	AGTTTTTCCG	GGACCGTAAA	ATATAATGGC	300
AGTAGCTATC CTTTCCCTAC	TACCAGCGAA	ACGCCGCGCG	TTGTTTATAA	TTCGAGAACG	360
GATAAGCCGT GGCCGGTGGC	GCTTTATTTG	ACGCCTGTGA	GCAGTGCGGG	GGGAGTGGCG	420
ATTAAAGCTG GCTCATTAAT	TGCCGTGCTT	ATTTTGCGAC	AGACCAACAA	CTATAACAGC	480
GATGATTTCC AGTTTGTGTG	GAATATTTAC	GCCAATAATG	ATGTGGTGGT	GCCCACTGGC	540
GGCTGTGATG TITCTGCTTG	TGATGTCACC	GTTACTTTGC	CGGACTACCC	TGGTTCAGTG	600
CCGATTCCTC TTACCGTTTA	TTGTGCGAAA	AGCCAAAACC	TGGGGTATTA	CCTCTCCGGC	660
ACACACGCAG ATGCGGGCAA	CTCGATTTTC	ACCAATACCG	CGTCGTTTTC	ACCTGCACAG	720
GGCGTCGGCG TACAGTTGAC	GCGCAACGGT	ACGATTATTC	CAGCGAATAA	CACGGTATCG	780
TTAGGAGCAG TAGGGACTTC	GGCGGTAAGT	CTGGGATTAA	CGGCAAATTA	CGCACGTACC	840
GGAGGGCAGG TGACTGCAGG	GAATGTGCAA	TCGATTATTG	GCGTGACTTT	TGTTTATCAA	900

(2) INFORMATION FOR SEQ ID NO:55:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 51 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:55:

GATCTGTTGA AGTTCCGGGT AGTCAGCATA TCGATAGTCA GAAAAAAGCT G

CLAIMS

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- 1. A method of targeting a bacterial adhesin to a specific location, comprising (i) identifying in said location adhesin-interacting receptor moiety which is recognizable by bacterial adhesins, (ii) isolating a bacterial cell that grows in said location and expresses an adhesin recognizing and interacting with said receptor moiety, and administering to the location the bacterial cell or the adhesin under conditions where the adhesin and the receptor moiety are brought into interacting contact whereby the adhesin is associated with the receptor moiety.
- 2. A method according to claim 1 wherein the receptor moiety is selected from the group consisting of a glycolipid, a glycoprotein, a protein, a polypeptide, a saccharide moiety and a peptide.
- A method according to claim 1 wherein the isolated bacterial cell expresses an adhesin having modified receptor moiety-binding properties relative to an adhesin natively expressed by the cell, the isolation of the cell comprising identifying in a parent bacterial cell, DNA sequence(s) coding for the binding domain(s) of said natively expressed adhesin and substituting at least one codon herein, whereby a modified adhesin molecule is expressed that is different in at least one amino acid from the adhesin expressed natively,
 and selecting a the bacterial cell expressing the modified adhesin having an altered adhesion phenotype relative to the natively expressed bacterial adhesin.
- 4. A method according to claim 1 wherein the bacterial cell expressing an adhesin that recognizes and binds to the receptor moiety is a recombinant bacterial cell derived from a parent bacterial cell that does not produce an adhesin binding to said receptor, by inserting into the parent cell a DNA sequence coding for an adhesin binding to the receptor

moiety, and selecting a bacterial cell expressing the DNA sequence.

- 5. A method according to claim 1 wherein a non-adhesin compound is associated with the adhesin, whereby said compound is targeted with the adhesin to the location comprising the receptor moiety recognizable by the adhesin.
- 6. A method according to claim 5 wherein the compound is covalently bound to the adhesin.
- 7. A method according to claim 6 wherein the compound is part of a fusion protein comprising the adhesin, the compound being selected from the group consisting of an enzyme, an antibody, an epitope and a toxin.
 - 8. A method according to claim 5 wherein the compound is one associated with the adhesin by a non-covalent binding.
- 9. A method according to claim 8 wherein the compound is selected from the group consisting of a pharmacologically active, a diagnostically active and an imaging compound.
 - 10. A method according to claim 1 wherein the specific location is a human or animal surface.
- 20 11. A method according to claim 1 wherein the specific location is a plant surface.
 - 12. A method according to claim 1 wherein the bacterial cell expresses a recombinant bacterial adhesin variant derived from a naturally occurring parent adhesin, said recombinant
- 25 bacterial adhesin variant having altered binding properties relative to the naturally occurring adhesin from which it is derived, the altered binding properties including binding to at least one receptor moiety to which the parent adhesin does not bind.

- 13. A method according to claim 12 wherein the adhesin variant is derived from a naturally occurring adhesin isolated from a cell structure selected from the group consisting of a capsule, a lipopolysaccharide layer, an outer membrane protein, a flagellum, a pilus, a fimbria, a non-fimbrial adhesin (NFA) and an afimbrial adhesin (AFA).
 - 14. A method according to claim 12 or 13 wherein the adhesin variant is a protein having an amino acid sequence differing in at least one amino acid residue from its parent protein adhesin.
 - 15. A method according to claim 14 wherein the adhesin variant is a FimH adhesin having an amino acid sequence which differs from the *E. coli* PC31 FimH adhesin as defined in Table 1 herein in at least one amino acid.
- 15 16. A method according to claim 15 wherein the FimH adhesin is one binding to a receptor selected from the group consisting of a domain where mannosyl residues are not terminal and a domain devoid of saccharide.
- 17. A method according to claim 15 wherein the adhesin vari-20 ant is a chimeric adhesin comprising amino acid sequences from different FimH adhesins.
 - 18. A method according to claim 15 wherein the FimH adhesin has an amino acid sequence which is selected from the group consisting of sequences appearing in Fig. 5 herein with designations CI#12, CI#4, CI#7 or CSH-50.
 - 19. A method according to claim 15 wherein the adhesin is one which, when tested for binding to yeast mannan (Mn), human plasma fibronectin (Fn), periodate treated Fn and the synthetic peptide FnSpl comprising the first 30 amino acids of Fn, only binds to Mn (M class).

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20. A method according to claim 15 wherein the adhesin is one which, when tested for binding to yeast mannan (Mn), human plasma fibronectin (Fn), periodate treated Fn and the synthetic peptide FnSp1 comprising the first 30 amino acids of Fn, binds to Mn and Fn (MF class).

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- 21. A method according to claim 15 wherein the adhesin is one which, when tested for binding to yeast mannan (Mn), human plasma fibronectin (Fn), periodate treated Fn and the synthetic peptide FnSp1 comprising the first 30 amino acid residues of Fn, binds to all of these (MFP class).
- 22. A method according to claim 15 wherein the adhesin is one which, when tested for binding to five Fn-fragments obtained by thermolysin treatment, only binds to the 40-kDa gelatin-binding fragment.
- 15 23. A method according to claim 22 wherein the adhesin is one which, when tested for binding to five Fn-fragments obtained by thermolysin treatment, binds to all five fragments.
 - 24. A method according to claim 15 wherein the adhesin is at least 90% homologous to the PC31 FimH adhesin.
- 20 25. A method according to claim 15 wherein the adhesin is a chimeric adhesin comprising amino acid sequences from different FimH adhesins.
 - 26. A method according to claim 15 comprising an amino acid sequence which differs from the *E. coli* PC31 FimH adhesin by at least one amino acid occurring between residues 27 and 119 of the mature FimH sequence.
- 27. A method according to claim 15 wherein the adhesin binds to a receptor moiety selected from the group consisting of a human receptor moiety, an animal receptor moiety, a plant receptor moiety and an inanimate, non-biological receptor moiety.

- 28. A method according to claim 1 wherein the bacterial cell being targeted is a cell comprising a gene coding for a gene product which, when expressed has a killing or cell function-limiting effect in said cell, the expression of said gene coding for the cell killing or cell function-limiting gene product being regulated in such a manner that the bacterial cell when targeted will be killed or limited in its function in a pre-determined manner.
- 29. A recombinant or mutant bacterial adhesin variant derived from a naturally occurring parent adhesin, said adhesin variant having altered binding properties relative to the naturally occurring adhesin from which it is derived, the altered binding properties including binding to at least one receptor to which the parent adhesin does not bind.
- 15 30. An adhesin variant according to claim 29 which is derived from a naturally occurring adhesin isolated from a cell structure selected from the group consisting of a capsule, a lipopolysaccharide layer, an outer membrane protein, a flagellum, a pilus, a fimbria, a non-fimbrial adhesin (NFA) 20 and an afimbrial adhesin (AFA).
 - 31. An adhesin variant according to claim 29 or 30 which is a protein having an amino acid sequence differing by at least one amino acid residue from its parent protein adhesin.
- 32. An adhesin variant according to claim 29 which is a FimH mannose-sensitive adhesin having an amino acid sequence which differs from the *E. coli* PC31 FimH adhesin as defined in Table 1 herein by at least one amino acid, said FimH adhesin binding to a receptor selected from the group consisting of a domain where mannosyl residues are not terminal and a domain devoid of saccharide.
 - 33. An adhesin variant according to claim 32 which is at least 90% homologous to the PC31 FimH adhesin.

- 34. An adhesin variant according to claim 32 which is a chimeric adhesin comprising amino acid sequences from different FimH adhesins.
- 35. An adhesin variant according to claim 29 which binds to a receptor moiety selected from the group consisting of an animal receptor moiety, a plant receptor moiety and an inanimate receptor moiety.
- 36. An adhesin variant according to claim 29 which is part of a fusion protein comprising the adhesin variant and a heterologous polypeptide.
 - 37 An adhesin variant according to claim 36 wherein the heterologous polypeptide is selected from the group consisting of an epitope, an enzyme, a toxic gene product and an antibody.
- 15 38. A FimH adhesin comprising 279 amino acids, having an amino acid sequence which differs from the *E. coli* PC31 FimH adhesin as defined in Table 1 herein by at least one amino acid.
- 39. A FimH adhesin according to claim 38 which has an amino acid sequence which is selected from the group of sequences appearing in Fig. 5 herein with designations CI#12, CI#4, CI#7 or CSH-50.
- 40. An adhesin according to claim 38 which binds to a receptor selected from the group consisting of a domain where 25 mannosyl residues are not terminal, a domain devoid of saccharide, a glycolipid, a glycoprotein, a protein, a polypeptide and a peptide.
- 41. An adhesin according to claim 38 which when tested for binding to yeast mannan (Mn), human plasma fibronectin (Fn), periodate treated Fn and the synthetic peptide FnSp1 compris-

ing the first 30 amino acids of Fn, only binds to Mn (M class).

- 42. An adhesin according to claim 38 which when tested for binding to yeast mannan (Mn), human plasma fibronectin (Fn), periodate treated Fn and the synthetic peptide FnSp1 comprising the first 30 amino acids of Fn, binds to Mn and Fn (MF class).
- 43. An adhesin according to claim 38 which when tested for binding to yeast mannan (Mn), human plasma fibronectin (Fn), periodate treated Fn and the synthetic peptide FnSp1 comprising the first 30 amino acid residues of Fn, binds to all of these (MFP class).
- 44. An adhesin according to claim 38 which when tested for binding to five Fn-fragments obtained by thermolysin treatment, only binds to the 40-kDa gelatin-binding fragment.
- 45. An adhesin according to claim 38 which when tested for binding to five Fn-fragments obtained by thermolysin treatment, binds to all five fragments.
 - 46. An adhesin according to claim 38 which is at least 90% homologous to the PC31 FimH adhesin.
- 47. An adhesin according to claim 38 which is a chimeric adhesin comprising amino acid sequences from different FimH adhesins.
- 48. An adhesin according to claim 38 comprising an amino acid sequence which differs from the *E. coli* PC31 FimH adhesin by at least one amino acid occurring between residues 27 and 119 of the mature FimH sequence.
 - 49. An adhesin according to claim 48 which binds to a receptor moiety selected from the group consisting of a human receptor moiety, an animal receptor moiety and a plant receptor moiety.

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- 50. A recombinant replicon comprising a DNA sequence selected from the group consisting of a sequence coding for a recombinant bacterial adhesin as defined in claim 29 and a sequence coding for a FimH adhesin as defined in claim 38.
- 5 51. A recombinant replicon according to claim 50 wherein the DNA sequence codes for a FimH adhesin having an amino acid sequence which differs from the *E. coli* PC31 FimH adhesin by at least one amino acid.
- 52. A replicon according to claim 52 in which the DNA sequence is at least 90% homologous to the PC31 fimH gene.
 - 53. A replicon according to claim 50 in which the DNA sequence is a chimeric fimH gene comprising DNA from different fimH genes.
- 54. A replicon according to claim 50 in which the DNA sequence comprises a DNA sequence which differs from the E. coli PC31 fimH gene by at least one codon between the codons coding for amino acid residues 27 and 119 of the mature FimH sequence.
- 55. A replicon according to claim 50 in which the DNA sequence comprises a further DNA sequence coding for a heterologous polypeptide.
 - 56. A replicon according to claim 55 wherein the polypeptide is selected from a group consisting of an epitope, an enzyme, a toxic gene product and an antibody.
- 57. A replicon according to claim 55 wherein the further DNA sequence codes for a gene product which is selected from a pesticidally active gene product and a pollutant-degrading gene product.
- 58. A replicon according to claim 50 wherein the DNA sequence is isolated from an *Enterobacteriaceae* species.

- 59. A fusion protein comprising an adhesin selected from the group consisting of a recombinant bacterial adhesin variant as defined in claim 29 and a FimH adhesin as defined in claim 38, and a heterologous polypeptide.
- 5 60. A fusion protein according to claim 59 wherein the heterologous polypeptide is selected from an epitope, an enzyme, a toxic gene product and an antibody.
 - 61. A fusion protein according to claim 59 which carries a non-covalently bound compound.
- 10 62. A bacterial cell which expresses an adhesin selected from the group consisting of a recombinant bacterial adhesin variant as defined in claim 29 and a FimH adhesin as defined in claim 38.
- 63. A recombinant bacterial cell according to claim 62 which comprises a recombinant replicon as defined in claim 50.
 - 64. A bacterial cell according to claim 62 which is selected from Enterobacteriaceae, Pseudomonadaceae, Vibrionaceae and Baccilaceae.
- 65. A bacterial cell according to claim 62 which further
 20 expresses a gene product selected from the group consisting
 of a pesticidally active compound, an immunologically active
 gene product and a pollutant-degrading active compound.
- 66. A bacterial cell according to claim 62 in which the recombinant adhesin variant is expressed as a fusion protein comprising the adhesin variant and a further polypeptide.
 - 67. A bacterial cell according to claim 62 which further comprises a gene coding for a gene product which, when expressed has a killing or cell function-limiting effect in said cell, the expression of said gene coding for the cell

killing or cell function-limiting gene product being regulated in such a manner that the bacterial cell when targeted to receptor in a specific location will be killed or limited in its function in a pre-determined manner.

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- 5 68. A method of isolating a bacterial cell expressing an adhesin having modified binding properties relative to a natively expressed adhesin, comprising identifying in the bacterial cell DNA sequence(s) coding for the binding domain(s) of said natively expressed adhesin and substituting at least one codon herein, whereby a modified adhesin molecule is expressed that is different in at least one amino acid from the adhesin expressed natively, and selecting a bacterial cell expressing the modified adhesin having an altered adhesion phenotype relative the natively expressed bacterial adhesin.
 - 69. A method according to claim 68 wherein a non-adhesin compound is associated with the adhesin.
- 70. A method according to claim 69 wherein the non-adhesin compound is associated with the adhesin by being expressed with the adhesin as part of a fusion protein comprising the adhesin.
 - 71. A method according to claim 68 which in a further step comprises binding non-covalently a compound to the adhesin when expressed.
- 25 72. A method according to claim 68 wherein the natively expressed adhesin is a FimH adhesin.
 - 73. A method according to claim 68 wherein the codon(s) is/are substituted by mutagenization.
- 74. A method of preparing a recombinant bacterial cell that
 30 binds to a specific receptor moiety, comprising introducing
 into a bacterium that does not produce an adhesin binding to

said receptor moiety, a DNA sequence coding for an adhesin binding to the receptor moiety, and selecting a bacterial cell expressing the DNA sequence.

- 75. A method according to claim 74 wherein the DNA sequence coding for an adhesin binding to the receptor moiety is a sequence coding for a FimH adhesin.
 - 76. A method according to claim 74 wherein the DNA is introduced by transforming the bacterial cell with a recombinant replicon as defined in claim 50.
- 10 77. A method according to claim 74 wherein a non-adhesin compound is associated with the adhesin.
- 78. A method according to claim 77 wherein the non-adhesin compound is associated with the adhesin by being expressed with the adhesin as part of a fusion protein comprising the adhesin.
 - 79. A method according to claim 74 which in a further step comprises binding non-covalently a compound to the adhesin when expressed.
- fimbriae which binds to a moiety to which the wild-type cell from which the mutant cell is derived does not bind, comprising contacting a population of said wild-type cell with said moiety, removing the contacted cells which do not bind to the moiety, cultivating cells binding to the moiety to obtain a culture which is enriched with regard to cells binding to the moiety and selecting from said culture a mutant cell binding to said moiety.
- 81. A method according to claim 80 wherein the moiety with which the wild-type cell population is contacted, is a casein.

- 82. A method of isolating a compound from a solution or suspension containing the compound, the method comprising contacting the solution or the suspension with a fusion protein according to claim 59 wherein the heterologous polypeptide has an affinity to the compound to be isolated.
- 83. A composition comprising a population of a bacterial cell as defined in claim 62.

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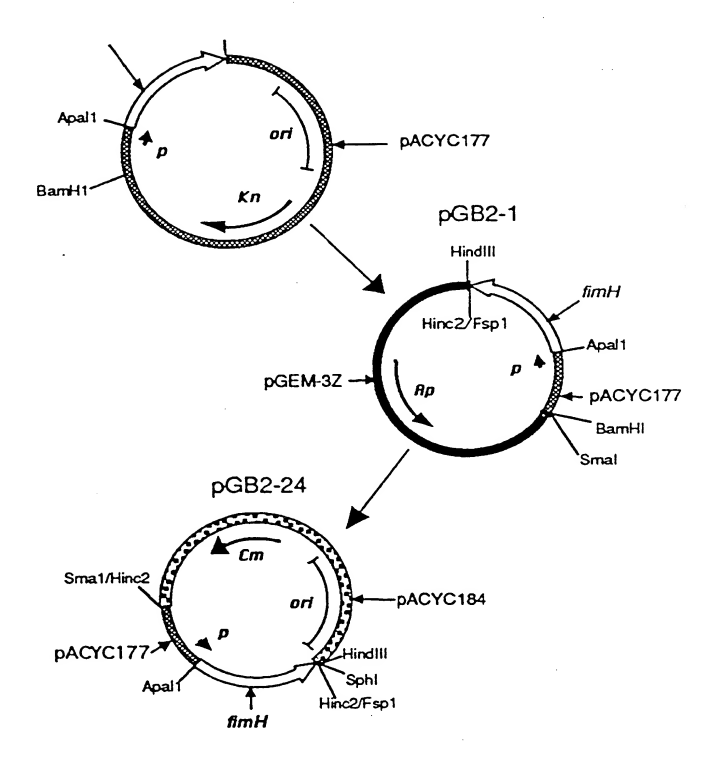


Fig. 1

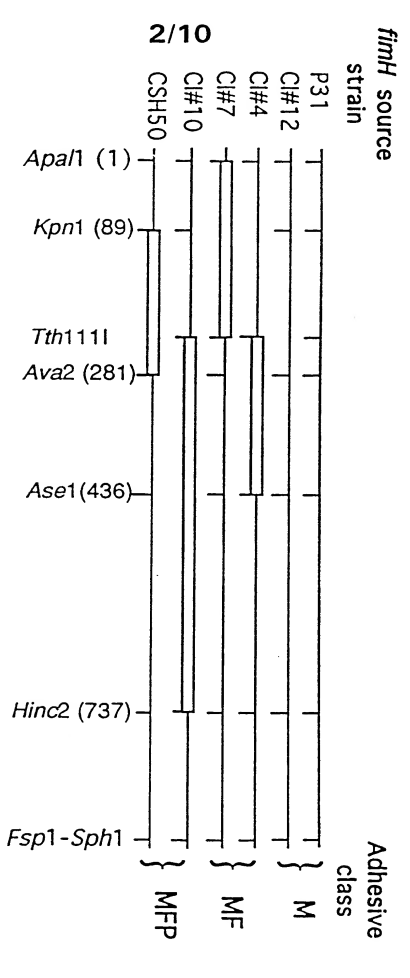
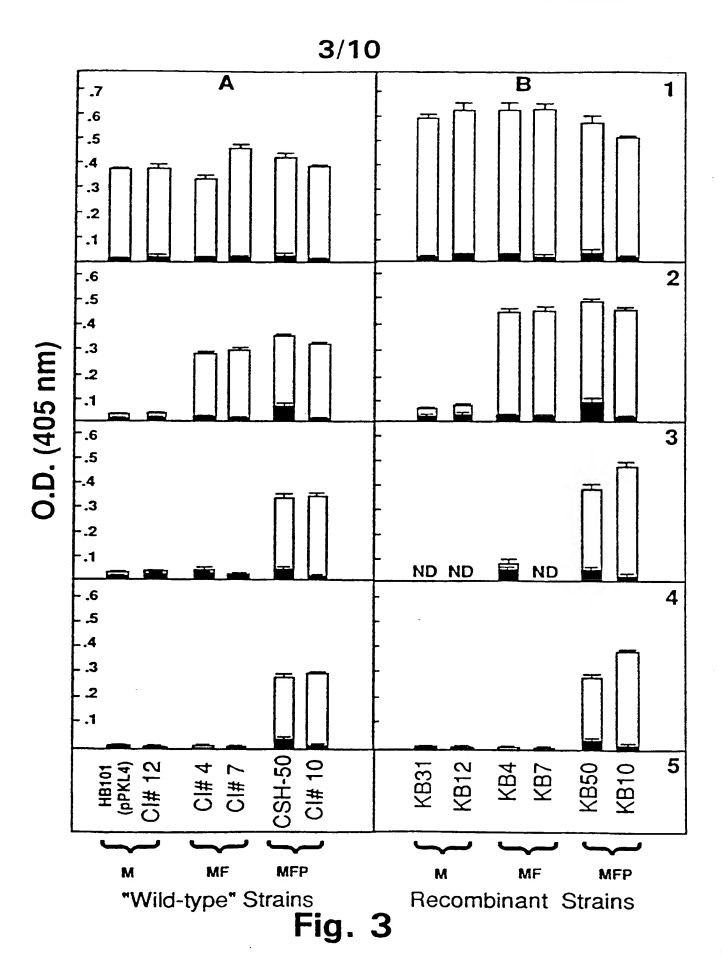


Fig. 2



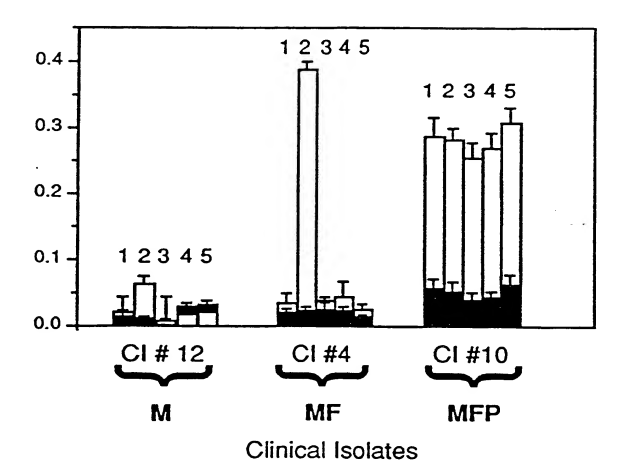


Fig. 4

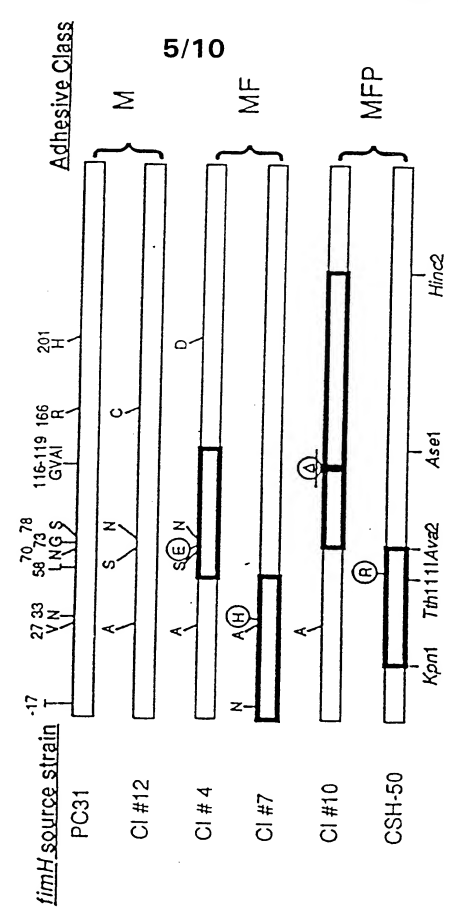


Fig. 5

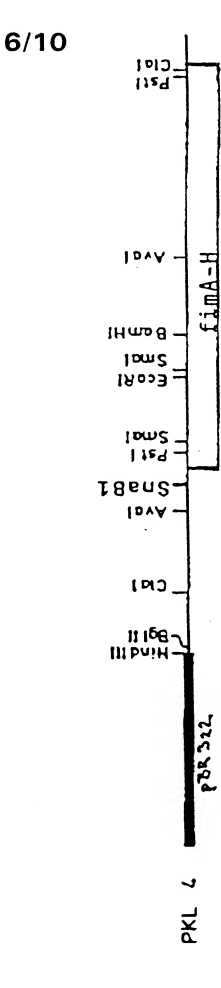


Fig. 6

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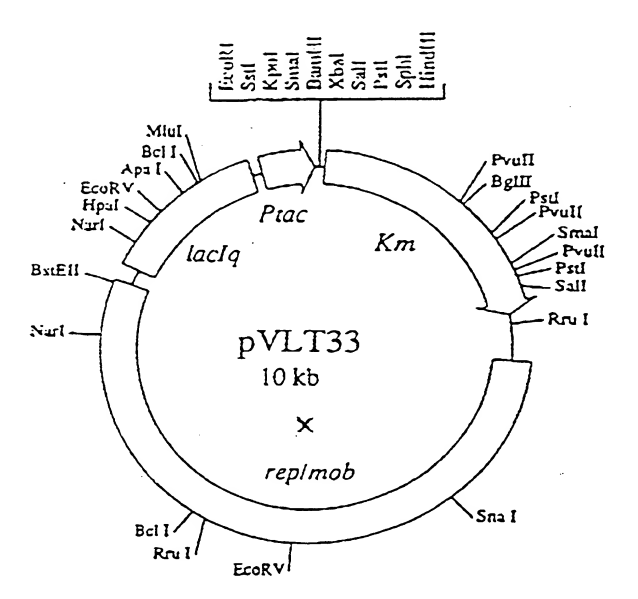


Fig. 7

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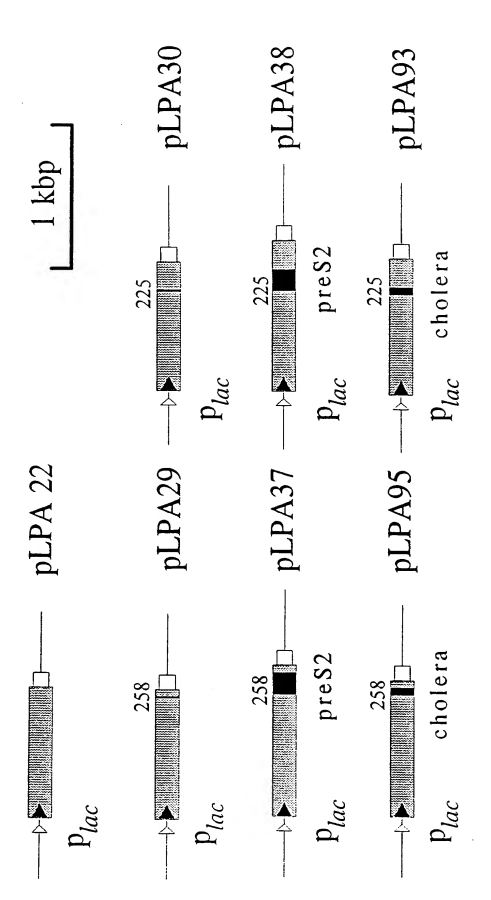


Fig. 8



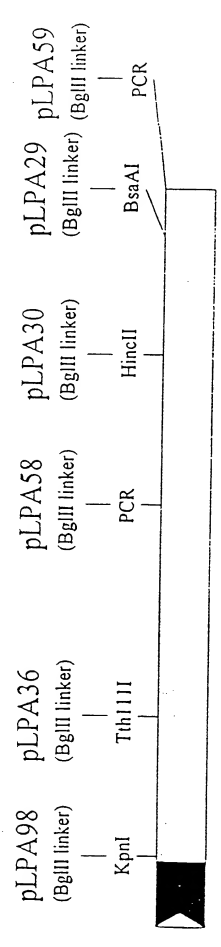


Fig. 9

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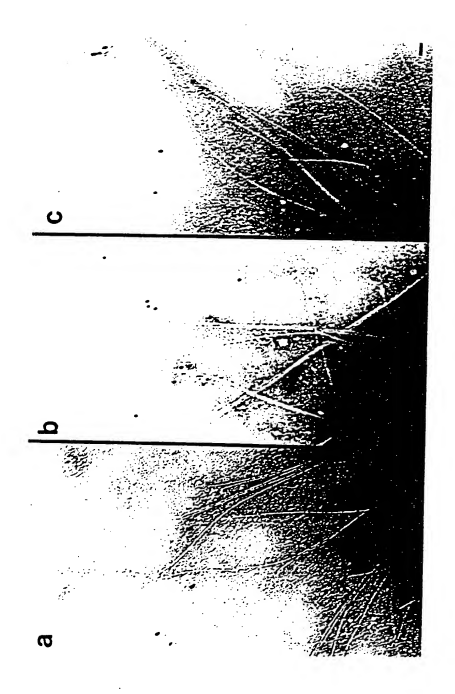


Fig. 10

A. CLASSIFICATION OF SUBJECT MATTER IPC 6 C12N15/31 C12N15/62

C12N1/21

C07K14/245

C12Q1/00

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) IPC 6

C07K C12N C12Q

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

C.	DOC	UM	ENT	5 (CON	SID	ERE	D	то	В	Εl	REL	EVA	NT
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Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	INFECTION AND IMMUNITY, vol. 60, no. 11, November 1992 WASHINGTON US, pages 4709-4719, EVGENI V. SOKURENKO ET AL. 'Functional heterogeneity of Type 1 fimbriae of Escherichia coli' see abstract see page 4709, right column, last paragraph - page 4710, left column, paragraph 1 see page 4712, right column, paragraph 1 see page 4714, right column, paragraph 1 see page 4715, right column, paragraph 1 page 4716, left column, paragraph 3 see page 4717, left column, paragraph 2	29,30, 38-40, 43,45, 62,64

Further documents are listed in the continuation of box C.	Patent family members are listed in annex.
Special categories of cited documents:	T' later document published after the international

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- mational filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
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- "&" document member of the same patent family

Date of the actual completion of the international search

Date of mailing of the international search report

26 June 1995

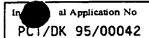
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016

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